MONOGRAPHS ON INORGANIC AND PHYSICAL CHEMISTRY

EDITED BY ALEXANDER FINDLAY, M.A., D.Sc., F.I.C.

THE ELECTRONIC CONCEPTION OF VALENCE AND THE CONSTITUTION OF BENZENE



THE

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WITH DIAGRAMS

LONGMANS, GREEN AND CO.

39 PATERNOSTER ROW, LONDON
FOURTH AVENUE & BOTH STREET, NEW YORK
BOMBAY, CALCUTTA, AND MADRAS

1921

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PREFACE.

THE electronic conception of valence as developed and applied in this monograph is concerned neither with the question of the ultimate nature of chemical affinity nor with the intimately related problem of the constitution of the atom. The variety of hypotheses now current, which deal with the constitution of the atom, has signally failed to furnish a uniform valence hypothesis which will enable chemists to cluculate chemical formulæ and reactions, or, in other words, to present more complete pictures of the relationships existing between the chemical constitution of substances and their chemical, physicochemical and physical properties.

After studying the many anomalous hypotheses on atomic structure and valence, the author has adopted the early and relatively simple suggestion of Sir J. J. Thomson that "if we interpret the 'bond' of the chemist as indicating a unit Faraday tube, connecting charged atoms in the molecule, the structural formulæ of the chemist can be at once translated into the electrical theory". Accordingly, the symbol—a short straight line between atoms—which indicates a "bond" in a structural formula, assumes an added significance since one end of the bond corresponds to a positive, the other to a negative charge, through the transference of an electron from the one atom to the other.

This, briefly, is the electronic conception of positive and negative valence which is employed herewith, solely as a formulative hypothesis. It is applied to the formulae of many substances but chiefly to the constitution of benzene.

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Methods are proposed and fully illustrated explaining how structural formula may be translated into electronic formula but only when it may be clearly demonstrated that the resultant electronic formula possess greater significance in interpreting and correlating chemical and physico-chemical phenomena than do our customary structural formula:

The monograph is divided into four parts. Part I. develops the electronic conception of positive and negative valence as a formulative hypothesis in chemistry. Part II. relates particularly to the constitution of benzene and its derivatives, and to the problem of substitution in the benzene nucleus. Part III. deals primarily with physical and physico-chemical properties, notably, molecular volumes, absorption of light and fluorescence. Part IV. considers the constitution of the metal-ammines and presents a bibliographical and chronological review of those articles of authors who have presented applications of the electronic conception of valence.

Concordant with the fact that the preface is usually the last part of a book to be written, a more comprehensive preface may be found in Section A of the final chapter.

In conclusion, I am deeply grateful to my colleague,

Dr. Earl F. Farnau, for valued suggestions, criticisms, and his reading of the manuscript, and also to Miss Eva Hauck, Secretary of the Department of Chemistry, University of Cincinnati, for the preparation of the index of names and the transcription of the manuscript.

University of Cincinnati, Cincinnati, Ohio, 18th May, 1920.

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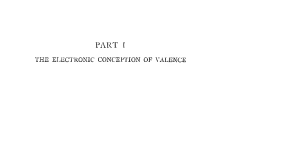
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CHAPTER I.

INTRODUCTORY.

THEORES of valency may be divided into two general classes. J. N. Firend states that "to the first of these belong such theories as those of Wemer and of Barlow and Pope, which postulate certain definite attractive or repellent forces, and then proceed to a discussion of the constitution of the molecule". These are designated as chemical theories of valency. Theories of the second class relate to the actual causes of chemical affinity and are influenced by the particular ideas of atomic structure held by their originators. Friend states that "the electronic theory of valency is a case in point, and as such is a subject for the physicist rather than the chemist."

Chemists, in general, will subscribe to this statement, because the various hypotheses on atomic structure and the fundamental nature of valence now in vogue are too diverse and too limited in their capacity either to interpret or to correlate definite chemical and physico-chemical phenomena. Be this as it may, there are certain simple aspects of the electronic conception of the constitution of the atom that may be translated directly into an electronic conception of valence. This, it will be shown, readily lends itself to the interpretation and correlation of many then to unexplained chemical and physico-chemical phenomena. In other words, the chemist may employ the electronic conception of valence as a formulative hypothesis in the field of chemistry.

The general purpose of this monograph is to show that the application of the electronic conception of positive and negative valences to the constituent atoms of elements and compounds leads to the development of a new type of structural chemical formula—the electronic formula—which is far more significant than the customary structural formula in its adaptation to the interpretation and correlation of chemical and physico-chemical phenomena.

To this end, there is presented a complete revision and an extended development of the subject matter of two senarate series of articles published by the author during the past twelve veaus One series, which appeared in the Zeitschrift fur physibalische Chemie under the general title Einige Anwendungen der Elektronischen Auffassung positiver und negativer Valenzen. presented new hypotheses on the relationship between chemical constitution and the phenomena of light absorption and fluorescence The other series appeared concurrently in the Journal of the American Chemical Society under the title Interpretations of Some Stereochemical Problems in Terms of the Electronic Conception of Positive and Negative Valences. In the majority of these articles an attempt was made to throw some light upon the manifold and ever recurrent problem of the constitution of benzene An application of the electronic conception of positive and negative valences to the constituent atoms of the benzene molecule culminated in the electronic formula of bensene. This formula is readily adaptable to and makes possible the interpretation of many chemical and stereochemical problems presented by benzene and its derivatives

My grateful acknowledgments are due to Si J J. Thomson whose earlier ideas on valency, as presented in his volume Electricity and Matter and subsequently elaborated in his Corpuscular Theory of Matter, directly led me to conceive, propose, and to apply the terms electroner and electronic tautomerism as fundamental concepts in the development of his electronic conception of valence. The electronic is a new type of isomer—the electronic isomer Electronic tautomewism involving electroners in dynamic countbrium.

The proposal of these new conceptions naturally involves the question of the actual existence of electroners. A few chemists have not, and others may not, be inclined to countenance the electronic conception of valence, electronic isomerism, and tautomeism chiefly upon the grounds that the independent existence of electronics has not as yet been fully established. In view of this attitude, it should be clearly understood that the question of the actual existence of electronics is a secondary matter which does not in any sense invalidate the use and sumificance of the electronic conception of velence as a formula-

two hypothesis in chemistry The conceptions of electronic isomerism and electronic tautomerism are necessary adjuncts of the electronic conception of valence Furthermore, these conceptions are necessarily demanded by and, therefore, warranted in the interpretation and correlation of many chemical phenomena. A statement of historic and prophetic interest, made by Kekulé in 1867, illustrates the principle at issue —

"The question whether atoms exist or not has but little significance from a chemical point of view its discussion belongs rather to metaphysics. In chemistry we have only to decide whether the assumption of atoms is an hypothesis adapted to the explanation of chemical phenomena. More especially have we to consider the question whether a further development of the atomic hypothesis promises to advance our knowledge of the mechanism of chemical observments.

"I have no hesitation in saving that, from a philosophical point of view. I do not believe in the actual existence of atoms. taking the word in its literal signification of indivisible particles of matter-I rather expect that we shall some day find for what we now call atoms a mathematico-mechanical explanation, which will render an account of atomic weight, of atomicity, and of numerous properties of the so-called atoms. As a chemist, however. I regard the assumption of atoms, not only as advisable, but as absolutely necessary in chemistry. I will even go further, and declare my belief that chemical atoms exist, provided the term be understood to denote those particles of matter which undergo no further division in chemical metamorphoses. Should the progress of science lead to a theory of the constitution of chemical atoms-important as such a knowledge might be for the general philosophy of matter-it would make but little alteration in chemistry itself. The chemical atoms will always remain the chemical unit; and for the specially chemical considerations we may always start from the constitution of atoms, and avail ourselves of the simplified expression thus obtained. that is to say, of the atomic hypothesis. We may, in fact, adopt the view of Dumas and of Faraday, that whether matter be atomic or not, thus much is certain, that granting it to be atomic, it would appear as it now does"

Apropos of the above quotation, let it be recalled that Kekulé's efforts to develop an understanding of the mechanism of chemical reactions are embodied to-day in the structure theory, the recognized foundation of the extensive achievements of Organic Chemistry, both theoretical and applied. The established utility of this structure theory, which is indeed a formulative hypothesis, is neither dependent upon the existence of atoms nor upon an intimate knowledge of the nature of valence or chemical affinity. Similarly, the utility of this electronic conception of positive and negative valences, also a formulative hypothesis with the structure theory as its foundation, is not primarily dependent upon the existence or the possible isolation of electronics.

The significance of this point of view may be more evident if some of the foregoing statements of Kekulé are paraphrased in modern terms: In chemistry we have to decide whether the electronic conception of valence is an hypothesis adapted to the explanation of chemical and physico-chemical phenomena More especially have we to determine whether or not the further development of electronic formulæ and the concentions of electronic isomerism and tautomerism promise to advance our knowledge of the mechanism of chemical reactions. In view of the fact that electronic formulæ, in many instances, have proven to be more precise and more significant than the customary structural formulæ in the explanation of physical and chemical phenomena and the mechanism of reactions, it is quite conceivable that the electronic conception of valence may become a necessary adjunct to the structure theory. This, of course, must depend upon the nature and extent of its applications and experimental verifications, and upon the part that should be played by just criticisms in bringing to light the relative meits and dements of its applications. The author hopes that these points of view will be constantly in the mind of the reader.

CHAPTER II

THE ELECTRONIC CONCEPTION OF VALENCE.

A. Fundamental Conceptions.

IN a lecture 1 on the Constitution of the Atom (Silliman Lectures, 1903), Sir J J. Thomson formulated the electronic conception of positive and negative valences in the following words.—

"On the view that the attraction between the atoms in a chemical compound is electrical in its origin, the ability of an element to enter into chemical combination depends upon its atom having the power of acquiling a charge of electricity. This, on the preceding view, implies either that the uncharged atom is unstable and has to lose one or more corpuscles before it can get into a steady state, or else that it is so stable that it can retain one or more additional corpuscles without any of the original corpuscles being driven out. If the range of stability is such that the atom, though stable when uncharged, becomes unstable when it receives an additional corpuscle, the atom will not be able to receive a charge either of positive or negative electricity, and will therefore not be able to enter into chemical combination. Such an atom would have the properties of the atoms of such elements as aroun or helium.

"The view that the forces which bind together the atoms in the molecules of chemical compounds are electrical in their origin, was first proposed by Berzelius; it was also the view of Davy and of Faraday Helmholtz, too, declared that the mightiest of chemical forces are electrical in their origin. Chemists in general seem, however, to have made little use of this idea, having apparently found the conception of 'bonds of ffinity' more fruitful. This doctune of bonds is, however, when regarded in one aspect almost identical with the electrical theory. The theory of bonds when represented graphically supposes that from each univalent atom a staught line (the symbol of a bond) proceeds; a bivalent atom is at the end of two such lines, a

tervalent atom at the end of three, and so on; and that when the chefnical compound is represented by a graphic formula in this way, each atom must be at the end of the proper number of the lines which represent the bonds. Now, on the electrical view of chemical combination, a univalent atom has one unit charge, if we take as our unit of charge the charge on the corpuscle: the atom is therefore the beginning or end of one unit Faradav tube the beginning if the charge on the atom is positive, the . end if the charge is negative A bivalent atom has two units of charge and therefore it is the origin or termination of two unit Faraday tubes Thus, if we interpret the 'bond' of the chemist as indicating a unit Faraday tube, connecting charged atoms in the molecule, the structural formulæ of the chemist can be at once translated into the electrical theory. There is, however, one point of difference which deserves a little consideration the symbol indicating a bond on the chemical theory is not regarded. as having duection, no difference is made on this theory between one end of a bond and the other. On the electrical theory, however, there is a difference between the ends, as one end corresponds to a positive, the other to a negative charge."

It is this earlier view of Su J. J. Thomson that is most readily and significantly adaptable to chemical formulae. The conceptions presented in the foregoing quotation may be illustrated and amplified by applying them to the combination of two univalent atoms, X and X, of such nature that X tends to lose a corpuscle, or election, which Y tends to acquire. Through the loss of one electron, i.e., one unit negative charge (represented

by the symbol \bigcirc), X functions positively; thus, $X - \bigcirc \rightarrow \overset{+}{\times}$ Through the acquisition of this electron, Y functions negatively;

Accordingly, the electronic formula of the resultant com- $\stackrel{\bullet}{\longrightarrow}$ pound, XY, from the union of $\stackrel{\star}{X}$ and $\stackrel{\star}{Y}$, is written $\stackrel{\star}{X}\stackrel{\longleftarrow}{\longrightarrow} Y$ which indicates the polarities of the bond of attraction or Faraday tube of force between X and Y.

In this connection it is of interest to note that Baly and Desch' consider the lable hydrogen atom, in keto-enol tauto-merism, to function as a potential ion "inasmuch as the bond of attraction or Paraday tube of force must be considered to be

lengthened sufficiently to allow of the interchange of the atom from the one postuon to the other within the molecule". They also extend this view to salts in solution maintaining "that the bonds of attraction connecting the 'rins' together are lengthened by the solvent. When the length of the Faraday tubes is below a certain critical length, the salt is 'non-ionized'. When the average length of the tubes of force is equal to or a little less than the critical length, a few interchanges of ions between adjacent molecules takes place, and the salt is partially ionized. When the length of the Faraday tubes is greater than the critical value, then perfectly free interchange takes place between thous of different molecules, and the salt is combetely 'clonized'."

From this point of view, the distinction between "electrolytes" and "non-electrolytes" is one of relation only; it depends upon the conditions which determine the critical lengths of the Faraday tubes between the constituent atoms of the substance.

Another phase of the electronic conception of valence has been presented by Sii William Ramsay⁹ in his Presidential Address to the Chemical Society (1908). He advanced the hypothesis that electrons are atoms of the chemical element, electricity; they possess mass, form compounds with other elements, and serve as the bonds of union between atom and atom. He employed this idea of valence to explain the extrusion of roitzable groups in such compounds as the cobaltammine intrites by further addition of ammonia. Such significant applications postend further development in the explanation of the mechanism of chemical changes.

A survey of these hypotheses leads to the conclusion, which is becoming more general, that, if the forces which hold the atoms together in electrolytes are electrical, then the same forces must also be assumed to hold in combination the atoms constituting the molecules of non-electrolytes. Hence, it may be maintained that chemical reactions which involve the dissociation of molecules, either of electrolytes or of non-electrolytes, are, let us say, electronic.

B. Electronic Formulæ of Diatomic Molecules.

Passing from this general conclusion, consider briefly some of the electrochemical conceptions and facts that are related directly to the determination of the electronic formulæ of the diatomic molecules of certain elements.

In the first place, it is of historical interest to note that one of the chief causes contributing to the overthrow of the Dualistic System of Berzelius 4 was his failure to apply the conception of electro-dualism to the constitution of elementary molecules. This, however, was not the case with Helmholtz,5 who, in his famous Faraday Lecture in 1881 applied an electrical conception of valence to the constitution not only of compound but also of elementary molecules These are his words: "If we conclude from the facts that every unit of affinity is charged with one equivalent either of positive or of negative electricity, they can form compounds, being electrically neutral only if every unit charged positively unites under the influence of a mighty electric attraction with another unit charged negatively. that even elementary substances, with few exceptions, have molecules composed of two atoms, make it probable that even in these cases electric neutralization is produced by the combination of two atoms, each charged with its full electric equivalent, not by neutralization of every single unit of affinity,"

This conception of Helmholtz is stated conversely, in the modern phraseology of ionic dissociation, by W. A. Noves, as follows. "If we suppose what seems not to be improbable, that all reactions involving the decomposition of molecules are preceded by an ionization of the parts of the molecules, it would follow that elementary molecules as well may ionize into positive and negative parts" Again, in this connection, Walden's researches on conductivities of substances other than acids, bases and salts in different ionizing media, prove that the halogens, bromine and iodine, furnish not only amons, but cations as well ++++ +++

Br and Br: I and L

Sir J. J. Thomson 8 states that this view "is also supported by the fact that when the molecules of an elementary gas are dissociated by heat, as in the case of rodine vapour, the electric conductivity of the dissociated gas is very high, showing that there are large quantities of both positive and negative ions present in the dissociated gas". Furthermore, "the numerous experiments which have been made on the dispersion of gases do not afford any evidence of the existence of any wide divergence between the dispersion of compound and elementary gases; hence we may conclude that if the atoms in the molecules of the compound gas are charged with electricity, the atoms in the molecules of elementary gases are also charged. Concurrently, the preceding facts indicate that the electronic formula and the electrolytic or electronic dissociation of a diatomic molecule (X₂) composed of univalent atoms may be represented by the following scheme.

$$X_n = X \xrightarrow{+-} X \rightleftharpoons X + X$$

C. Electronic Isomers or Electromers.

An extension of these ideas to the chemical union of two elementary gases, X_2 and Y_3 , develops some new and fundamental conceptions. Ordinarily, such a reaction is represented by the equation $X_2 + Y_2 \rightleftharpoons 2XY$, but in terms of electronic formulas, combination, preceded by electronic dissociation, is represented by the following scheme-

Ordinarily only one structural formula is assigned to the compound XY, namely, X——Y; but, in the above scheme it is evident that XY may be represented by two different electronic

formulæ, namely, X—Y and X—Y. I have proposed the term electronic isomer, or more briefly, electronics of this new type of isomer. Several different electronic formulæ may be attributed to a given structural formulæ but this does not necessarily imply that the several electronic isomers or electromers are capable of independent existence. Sin J. J. Thomson "foresaw the possibility of this new type of isomerism, and stated that even if such isomers were stable they would possess very different degrees of stablity, and "it must be remembered that in considering the possibility of the existence of isomers from purely geometrical considerations, all questions as to stablity are ignored, so that isomers which are indicated by geometry as possible may be dynamically unstable and thus incapable of

proparation". Detailed evidence for the existence of electromers of the types $X \stackrel{-}{---} Y$ and $X \stackrel{-}{---} Y$ will be presented in subsequent chapters.

D. Electronic Tautomerism.

The dynamic instability referred to above serves as the basis of another new conception, namely, that of electronic tautomerism. In the usual sense, signifies an equilibrium mixture of ordinary structural isomers as, for instance, the well-known lactonic and enolic modifications. Electronic tautomerism signifies an equilibrium mixture of two (or more) electronies in the sense that one electronier may be assumed to revert to another electromer through the thansposition of valence electronism. This will effect a reveisal of the polarity of certain atoms (or tadicals) in the respective electromers as indicated in the following detailed scheme:

$$x \xrightarrow{+-} Y \rightleftharpoons x + y \rightleftharpoons x + \Theta + Y \rightleftharpoons x + Y \rightleftharpoons x \xrightarrow{-+} Y$$

It should be noted that in the transition from electromer $\overset{+}{X}.\overset{-}{Y}$ to $\overset{-}{X}.\overset{+}{Y}$, or vice verse, abbreviated thus

$$(X \xrightarrow{+} \overline{-} Y \rightleftharpoons X \xrightarrow{-} \overline{+} Y),$$

when \vec{Y} loses an electron, it becomes a neutral atom or radical; and \vec{X} , acquiring this electron, or unit negative charge, also becomes a neutral atom or radical. The intermediate existence of neutral atoms or radicals. The antermediate existence and transparent electronic tautomerism, furnishes a fundamental point of view for the interpretation of the existence and properties of "free radicals" (notably triphenyl-methyl and its derivatives) and the so-called "inascent state". These features will be developed in subsequent chapters. One of the pulposes of this monograph is to show that many chemical reactions and their interpretation fully substantiate the principle of electronic tautomerism or the existence of electroners in dvanuer caulibilium.

E. The Electronic Valence of an Atom.

It has been noted that a neutral atom (the valence in the neutral state is zero) may acquire one unit positive charge through the loss of one electron, or it may acquire one unit negative charge through the acquisition of one electron. Such a nuivalent atom may function, either as an independent ion or in chemical combination with another atom, in two distinct ways eccording as it is positively or negatively nuivalent. Thus, a maintain thydrogen atom may function as H or as H; and the

univalent hydrogen atom may function as H or as H; and the univalent chlorine atom as Cl or as Cl.

Through (1) the gain or (2) the loss of two electrons, or (3) through the simultaneous loss of one electron and the gain of another electron, a bivalent atom may function in biree distinct ways. Accordingly, its valences are respectively (1) both hegative, or (2) both positive, or (3) one positive and one negative. Thus, a bivalent oxygen atom presents biree types:

O, O, O. Similarly, the *tervalent* nitrogen atom presents four types: N, N, N, and N. The *quadrivalent* carbon atom functions in five ways:—

It thus becomes evident that an atom whose valence is (n) may function electronically in (n+1) different ways. This rule 12 is used in figuring the number of electronic formula which may be attributed to a given structural formula. Its application to the derivation of the electronic formula of benzene and other compounds is presented in detail later. In the meantime, it is expedient to illustrate this rule by a number of simple reactions and electronic formula, and, simultaneously, to consider the methods commonly employed in deliving the electronic formulae of elements and compounds

CHAPTER III.

METHODS OF DEVELOPING ELECTRONIC FORMULÆ.

It has been noted that electronic formulae are virtually structural formulæ in which the bonds are qualified as having direction in the sense that one end of a bond corresponds to a positive, the other end to a negative charge. The structural formula of a substance is derived through the consideration of three factors (1) the number of atoms in the molecule; (2) the valence of each atom, and (3) the chemical and physical properties of the substance in question. As stated by Perkin and Kipping. 13 of the state of combination of all the atoms of which the molecule is composed, and to express the results in a structural formula, this formula then not only shows the constitution or structure of the compound, but also summarizes in a concise and simple manner the more important chemical properties of the compound.

Now the conversion of a structural formula into an electronic formula by assigning positive and negative chaiges to the opposite ends of the bonds is not an arbitrary procedure. it must be governed by a careful study of certain phenomena, notably (A) Ionization and Electrolysis, (B) Oxidation-reduction processes, and (C) Hydrolytic teactions. Each of these phenomena may be interpreted very aptly in terms of electric charges, or, in other words, in terms of positive and negative valentes: The correlation of these phenomena and principles makes the development of an electronic formula a consistent procedure. Simple illustrations of a general type may now be considered.

A. Ionization and Electrolysis.

In an aqueous solution, hydrogen chloride molecules by dissociation yield positive hydrogen and negative chlorine ions. The former migrate to the negative cathode while the latter

move to the positive anode. Hence the dissociating molecules of hydrogen chloride are qualified by the electronic formula H--Cl, and their electrolytic dissociation by the scheme $H \stackrel{\top}{-} C1 \Rightarrow H + C1.$

This view does not preclude the possible existence, even in an aqueous solution, of negative hydrogen and positive chlorine ions The following illustration will show how, during electrolysis, negative hydrogen ions may be assumed to result by cathodic reduction, and positive chlorine ions arise by anodic oxidation, a hydrogen atom carrying a unit positive charge may

lose it on contact with the cathode; H + O -> H. The resultant neutral atom immediately acquiring an electron from the cathode

becomes a negative hydrogen ion, H + ⊖ → H, which is naturadly repelled from the cathode and immediately combines with an approaching positive hydrogen ion to form a molecule of

hydrogen,
$$\ddot{H} + \ddot{H} \rightarrow \ddot{H} - \ddot{H} = H_2$$

An analogous and simultaneous process may take place at the anode, in which case the negative chlorine ion gives up one electron to the anode thereby becoming a neutral atom, which through the loss of another election becomes a positive chlorine atom. The union of a positive and a negative chlorine ion vields the molecule Cl.----Cl.

$$\overset{-}{\text{Cl}} - \overset{-}{\Theta} \Rightarrow \overset{-}{\text{Cl}}, \overset{-}{\text{Cl}} - \overset{+}{\Theta} \Rightarrow \overset{+}{\text{Cl}};$$

$$\overset{+}{\text{Cl}} + \overset{-}{\text{Cl}} \Rightarrow \overset{-}{\text{Cl}} \xrightarrow{-\text{Cl}} = \overset{-}{\text{Cl}}_{*}.$$

This scheme leads to the definition of oxidation and reduction in terms of electrical charges or electrons.

R Oxidation and Reduction.

Anodic oxidation and cathodic reduction are effected by means of the electric current. Electric currents may be produced by means of oxidation and reduction reactions. Hence oxidation is generally considered to involve the acquisition of positive, or the loss of negative electrical charges, by atoms or ions; and, reduction involves the acquisition of negative, or the loss of positive charges. Now lince the electron is the unit negative charge 16

of electricity, equivalent to the charge on a univalent ion, oxidation may be defined, in terms of the electronic conception, as the loss of electrons by atoms or ions, while reduction is the gain of electrons by atoms or ions.

The deplacement of hydrogen from an acid by means of a metal is represented by an equation of the following type: M (a bivalent metal) + $2HX \rightarrow MX_2 + H_2$. The ronic equation for this reaction is $M + 2H \rightarrow M + H_2$. The metal, M, is oxidized to the bivalent ion, M, while the hydrogen rons, 2H, are reduced to molecular hydrogen, H. A more detailed analysis of this change is presented in terms of the electronic conception according to the following scheme:—

$$\begin{array}{llll} \mathbf{M} &=& \stackrel{+}{\mathbf{M}} \stackrel{-}{\longrightarrow} \stackrel{+}{\mathbf{M}} \stackrel{+}{+} & : \bigodot, \\ \\ \stackrel{+}{\mathbf{H}} &:& \bigodot \longrightarrow \mathbf{H}, \ \mathbf{H} &+& \bigodot \longrightarrow \stackrel{-}{\mathbf{H}}, \\ \\ \stackrel{+}{\mathbf{H}} &+& \stackrel{-}{\mathbf{H}} \longrightarrow \mathbf{H} \stackrel{+}{\longleftarrow} \mathbf{H} &=& \mathbf{H}_{2} \end{array}$$

By way of explanation, an atom of the metal, M, is oxidized to the bivalent ion \widetilde{M} through the loss of two electrons, one of which reduces $\overset{\circ}{H}$, first, to neutral hydrogen, H. This neutral atom is then further reduced by the other electron to $\overset{\circ}{H}$. The union of $\overset{\circ}{H}$ and $\overset{\circ}{H}$ yields molecular hydrogen. This process closely resembles the *cathodic reduction* of positive hydrogen ions during the electrolysis of aqueous solutions of acids as previously noted

These illustrations show that the number and the polarity of the charges of a given atom indicate its state of oxidation or reduction. Conversely, the state of oxidation of a given atom in a compound may be readily correlated with the electronic formula of the compound. The significance of this punciple becomes more evident if attention is directed to the polarities of the valences of the carbon atom in five typical compounds, namely, (I.) methane, (II.) methyl alcohol, (III) formaldehyde, (IV.) formic acid, (V) carbonic acid and its anhydride, carbon dioxide. If, in these compounds, the hydrogen actors function positively

and the oxygen atoms are negatively bivalent, then the electronic formulæ of compounds I.-V. inclusive are written as follows.-

A rule has been stated, namely, that when the valence of an atom equals (n), that atom may function electronically, or in electronic formulæ, in (n + 1) different ways. Since the carbon atom is quadrivalent, it functions in five different ways, thus—

These types of carbon atoms (I.-V) are embodied, respectively, in the preceding electronic formulæ of (I,) methane, (II.) methyl alcohol. (III) formaldehyde. (IV.) formic acid, and (V) carbonic acid. Now since the acquirement of electrons or negative valences corresponds to reduction, and the loss of electrons, i.e., the development of positive valences, corresponds to oxidation, it is at once evident that the above types and electronic formulæ (I -V. inclusive) represent the successive stages of oxidation of the carbon atoms in the transitions from methane to carbon dioxide. In fact Bone has demonstrated that the slow combustion (i.e., oxidation) of the hydrocarbon methane at temperatures below its ignition point, may be regarded as involving the successive formation of methyl alcohol, formaldehyde, formic acid, and finally, carbonic acid or carbon dioxide. The electronic formulæ of these compounds afford an interpretation of these oxidations in terms of the positive and negative valences of the constituent atoms. Such relationships as those noted in the preceding types (I.-V) are made manifest, not by the ordinary structural formulæ, but only by the electronic formulæ of these five typical carbon compounds. In this way, the extensive field of exidation-reduction reactions not only affords an experimental justification for the use of the electronic conception of positive and negative vulences but also serves as one method of deriving electronic formula.

C. Hydrolytic Reactions.

Another method of developing electronic formulæ and of correlating them with chemical properties is found in an electronic interpretation of hydrolytic reactions.

Granting that the hydrogen ion is positive and that the hydroxyl ion is negative under normal conditions, i.e., provided the hydrogen ion is not reduced and that the hydroxyl ion is not oxidized, and that, accordingly, the electronic formula of

water is H—O—H, then it follows that hydrolytic reations afford an experimental method for ascertaining the polarity of the radicals of a compound under the particular conditions of the hydrolysis.

The applicability of such a method both to electrolytes and to non-electrolytes is indicated by Abegg. "The observation that all reactions in which ions participate in measurable amounts—even the hydrolytic actions of the extremely weakly dissociated water—proceed to their equilibra with an immeasurably great velocity, has induced the assumption that indeed every capacity to react is to be attributed to the presence of ions. A basis for this assumption has been thought to exist in the fact that reactions between non-electrolytes usually proceed with extreme slowness corresponding to an immeasurably small, but not absolutely non-existent, dissociation." The method of applying hydrolytic factions to the determination of the polarity of the radicals of either electrolytes or of non-electrolytes may be illustrated in general as follows.

If a compound, X.Y, on hydrolysis yields compounds of the types H X and HO.Y, then X.Y is qualified by the electronic

formula X — Y. Its hydrolysis proceeds according to the scheme:—

formula X — Y, the hydrolysis of which proceeds according to the abbreviated scheme:—

Thus, the products of hydrolysis of X.Y indicate whether its electronic formula is $\bar{X} \stackrel{+}{Y} = \bar{X} \stackrel{+}{X} = \bar{X}$.

In some instances, to be considered in detail later, a given compound, X Y, will yield four different products on hydrolysis, namely, H,Y, HO.X, H.X, and HO.Y. In such instances, the conclusion is evident that the compound X.Y must have

functioned in two distinct ways, namely, as $\overset{+}{X}.\overset{-}{Y}$ and $\overset{-}{X}.\overset{+}{Y}$; or, $\overset{-}{i}$, in other words, X.Y presents an example of electronic tauto-

$$(\overset{\downarrow}{x}\overset{\downarrow}{y}\overset{\downarrow}{\Rightarrow}\overset{\downarrow}{x}\overset{\downarrow}{y}) + \overset{\downarrow}{u}\overset{\downarrow}{o}H$$

This chapte has endeavoued to show that the development of an electronic formula, or the translation of a structural formula into an electronic formula, is made possible through the study and interpretation, in terms of positive and negative valences, of plenomena of ionization, oxidation-eduction processes, and hydrolytic reactions. The principles thus outlined will be illustrated fully in subsequent chapters with the purpose of making more appaient the enhanced significance of electronic formulæ.

CHAPTER IV.

RLECTRONIC AMPHOTERISM.

THE first rule relating to the positive and negative valencies of an element was proposed by Abegg and Bodlander 14 and subsequently by Abegg. 15 These authors developed a theory of valency—noimal and contravalency—of opposite polarity. The former is the stronger and contesponds, as its name implies, to the accepted valencies of the element. The normal valencies are usually positive in metals but negative in non-metals and the numerical value is equal to eight minus the number of contravalencies. The Periodic Classification of the elements furnished the basis for the distribution of the normal and contravalencies.

A few years later Friend ¹⁶ developed an electronic theory of positive and negative valence in which he distinguished between (1) free negative valency, (2) fiee positive valency, and (3) residual or latent valency.

Friend defines the fice negative valence of an element in terms of its capacity to combine with hydrogen: "Since hydrogen is electro-positive and monovalent, only those elements which possess a negative valence can combine with it. As Ramsay has pointed out this is equivalent to saying that only those elements which are capable of receiving electrons can form hydrides". In addition to negative, some elements possess positive valency, that is to say, they are also able to part with electrons. Such elements may be termed amphoteric. Others apparently possess only nositive or only negative valency.

Referring to free positive valency, Friend states that "the numerical value of the positive valency is not so easily determined as that of the negative, since we have no negative element corresponding with hydrogen, which combines with one atom of any other element in only one proportion". However, Friend

suggests fluorine as the most suitable element for determining the maximum positive valency of the elements and maintains that "chlorine, fluorine, and oxygen are the only elements which possess free negative but no free positive valencies".

Finally, Friend's conception of residual or latent valency is quite similar to the "neutral affinites" of Spiegel and the leath toil double vollencies of Arthenus, in that latent valences can be called out in pairs of equal and opposite sign. Thus, "In Ramsay's phraseology, this is equivalent to saying that when an element exerts its latent valency, it smultaneously parts with and receives, an electron. Consequently, the sum total of the electrons remains the same, and the electrochemical properties of the atom are unaltered!

I am constrained to maintain that the foregoing rules of positive and negative valencies as proposed by Abegg and Bodlander, and by Friend, may be replaced by a more general and more comprehensive rule, namely, that when the given valency of an element equals n, that element may function in (n+1) different ways. This rule has been illustrated in the case of the quadrivalent carbon atom and its typical compounds. In terms of this rule the univalent hydrogen atom may function in two ways, i.e., either positively or negatively. Accordingly, hydrogen in hydrogen chloride is positive and the

electronic formula of the hydrogen chloride molecule is H — Cl; but, in sodium hydride, if sodium is positive, hydrogen must be negative, thus Na⁺—H. This is in direct opposition to Friend's statement that "since hydrogen is electropositive and monovalent, only those elements which possess a negative valence can combine with it".

In like manner, univalent chlorine may function in two ways: it is negative in hydrogen chloride but positive, as will be shown later, in hypochlorous acid, H⁺—C⁻—Cl, and other compounds. This, too, is in opposition to Friend's statement ¹⁸ that "chlorine, fluorine, and oxygen are the only elements which

possess free negative but no fiee positive valencies". Chlorine, at least, possesses positive valence as will be shown subsequently.

The capacity of a given element to lose and to acquire electrons and thereby function in various ways may be termed

electronic amphoterium. It is now essential to subsequent developments to consider some instances in which the electronic amphoterism of hydrogen and chlorine is manifested. Since it is generally conceded that hydrogen normally functions positively while chlorine normally functions negatively, it will be of more particular interest to discuss the reactions and the electronic formulæ of a few simple compounds in which hydrogen is negative, and others in which chlorine is positive.

A. Negative Hydrogen.

The conception of negative hydrogen is not new. Brodle. ³ in 1850, assumed the existence of a relation (polarity) between atoms entering into combination of such kind that one atom is distinguished as positive or negative as contrasted with the other. He represented the evolution of hydrogen which occurs when copper hydride is treated with hydrochloric acid, in terms of the ded equivalents, thus—

This is parallel to our present assumption that both copper hydride and molecular hydrogen may contain hydrogen atoms which function negatively.

It has been noted previously that negative hydrogen may result through cathodic reduction of positive hydrogen ions in the electrolysis of aqueous solutions of acids and precede the formation of molecular hydrogen. Likewise, it may be formed and precede the liberation of hydrogen when metals interact with acids or with water. Direct evidence of the existence of negative hydrogen is found in certain reactions of sodium hydride, and reference should be made to the interaction of sodium, on the one hand, and of sodium hydride on the other, with water according to equations (1) and (2), respectively:—

From an ionic standpoint these reactions are represented thus .--

(1)
$$zNa + zH \rightarrow zNa + II_0$$

(2) $NaH + H \rightarrow Na + H_0$

A further analysis of these reactions from an electronic point of view requires that a molecule of metallic sodium be regarded simply as a compound of a positive sodium ion with a negative electron, the interaction of which with water (i.e., with the positive hydogen ions) corresponds to the following scheme:—

$$2NA = 2\stackrel{\dagger}{NA} \bigcirc \rightleftharpoons 2\stackrel{\dagger}{NA} + 2 \bigcirc$$

$$\stackrel{\dagger}{\Pi} + \bigcirc \Rightarrow \Pi, H + \bigcirc \Rightarrow \stackrel{\dagger}{\Pi},$$

$$\stackrel{\dagger}{H} + \stackrel{\dagger}{\Pi} \Rightarrow \stackrel{\dagger}{H} \stackrel{-}{\longrightarrow} H = H_0.$$

In other words, metallic sodium is oxidized to ionic sodium through the loss of electrons which reduce ionic hydrogen to the molecular state. Analogously, on the other hand, sodium hydride is a compound of positive sodium and negative hydrogen

atoms, NaH Hence, the ionic equation (2) above is represented electronically as follows —

$$Na.H = NaH \rightleftharpoons Na+H$$
 $+ H \Rightarrow H.H = H.$

In other words, sodium hydride may be regarded as an ionogen which, on dissociation, yields positive sodium and negative hydrogen ions. The latter combine with positive hydrogen ions yielding molecular hydrogen.

Further evidence that the hydrogen of sodium hydride functions negatively is shown by the formation of sodium formate according to the equation—19

$$Na.H + CO_g \rightarrow HCO_gNa.$$

An interpretation of this reaction from the electionic point of view shows it to be an oxidation-reduction process. It has been noted previously that each of the four valencies of the carbon atom in carbon dioxide is positive while in formic acid one of the carbon valences is negative and the other three are positive. It follows, therefore, that when sodium hydride interacts with carbon dioxide, one of the four positive valences of the carbon atom is reduced to a negative valence through the reducing action of the negative hydrogen atom of sodium

^{*} See Electronic formulæ on p. 17

hydride. A detailed analysis of these changes may be indicated schematically as follows:---

(a) Na. II
$$\Rightarrow$$
 Na + \hat{H} .
(b) $\hat{H} \rightarrow \hat{H} + 2 \ominus$
(1) CO₂ = $0 \xrightarrow{-+} \stackrel{+}{+} \stackrel{-}{-} 0 \Rightarrow 0 \xrightarrow{-+} \stackrel{+}{+} \stackrel{-}{-} 0$
(a) $0 \xrightarrow{-+} \stackrel{+}{+} \stackrel{-}{-} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{-+} \stackrel{+}{+} \stackrel{-}{-} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{-+} \stackrel{+}{+} 0 \xrightarrow{--} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{-+} \stackrel{+}{+} 0 \xrightarrow{--} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{-+} 0 \xrightarrow{--} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{-+} 0 \xrightarrow{--} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \longrightarrow + 2 \ominus \Rightarrow 0 \xrightarrow{--} 0 \longrightarrow + 2 \ominus \Rightarrow 0 \longrightarrow + 2 \bigcirc \to 0 \longrightarrow + 2 \longrightarrow + 2 \bigcirc \to 0 \longrightarrow + 2 \bigcirc \to$

In (1), dissociation of NaH yields Na and H, In (2), the unstable H becomes H through the loss of two electrons, 20, and thereby acts as a reducing agent. Note that (3) illustrates the "opening up" of one of the double bonds of carbon dioxide presenting one free negative and one free positive valence, so to speak. In (4), the electrons, 20, liberated in the transition of H to H, effect the reduction of the free positive valence of carbon dioxide to a free negative valence. In (5), the union of Na from (I), and of H from (2), with the respective free negative valences

There is another scheme of interpreting the reaction in question. The dissociation of sodium hydride according to (1) into Na and H, in conjunction with the opening up of the double bond in carbon dioxide, as noted in (3), makes possible the duect

of the oxygen and carbon atoms of carbon dioxide from (4), completes the electronic formula of sodium formate.

additions of Na and H to the free negative and the free positive valence, respectively, of carbon dioxide, thus:-

The instability of \widetilde{H} and its tendency to revert to \widetilde{H} $(\widetilde{H} \to \widetilde{H} + 2 \bigoplus)$ with the loss of two elections which reduce the positive carbon valence to a negative valence, is the occasion of the translable electioner, containing negative

hydrogen,
$$O \xrightarrow{-+} C \xrightarrow{+-} O \xrightarrow{-+} Na$$
, to the stable electromer,

that in the change from the unstable electromer (NaO_2C^-H) to the stable electromer (NaO_2C^-H) , the \bar{H} of the former has been oxidized to \bar{H} in the latter, and, simultaneously, the positive carbon valence of the former electromer has been reduced to a negative valence in the latter. Hence this reversal of the polanty of the carbon-hydrogen valence affords an example of an electrome intranolecular axidation-eduction \bar{I} thould here be emphasized that any electronic intramolecular oxidation-reduction reaction virtually corresponds to the transi-

tion of one electromer to another $(X \xrightarrow{-} Y \rightleftharpoons X \xrightarrow{-} Y)$ previously noted (p. 12) and defined as electronic tautomensm. Whether one electromer, or the other, or both, are the products of a given reaction will depend upon certain conditions which will be considered later.

Naturally it would be expected that the hydrides of stongly electropositive metals, such as sodium, contain negative hydrogen, as has been shown. It is interesting and significant to note that the hydride of the non-metal silicon also functions as a compound

of negative hydrogen. This is attested by the interaction (hydrolysis) of silicon hydride and aqueous potassium hydroxide. One volume of the former yields in alkaline solution four volumes of hydrogen according to the equation—

Now, potassium silicate is analogous in composition to potassium carbonate, a derivative of carbonic acid in which each of the valences of the carbon atom is positive. Accordingly, each of the valences of the silicon atom in potassium silicate is assumed to be positive. Furthermore, since neither water no potassium hydroxide acts as a reducing agent in the above equation, it must also be concluded that each of the valences of the silicon atom in silicon hydride is positive. Hence the hydrogen atoms of silicon hydride function negatively and the above section from the electronic standopoint is represented as follows:—

In this connection it is worthy of note that silicon and carbon, from the standpoint of the periodic classification, belong to the same natural family. Since the non-metal silicon may be united to negative hydrogen as shown above, it is natural to conclude that carbon atoms also may hold in combination megatively functioning hydrogen atoms. This condition will be met in the electronic formula of benzene. That thee of the hydrogen atoms of the benzene molecule, \mathbb{Q}_2H_n , function negatively while the other three function positively will be considered fully in subsection that parts.

In analogy with the hydrolysis of silicon hydride, silicon tetrachloride is also completely hydrolyzed in accordance with the scheme:—

$$\begin{array}{c} Cl & 0^{-\frac{1}{2}H} \\ + Cl & + H^{\frac{1}{2}-0-\frac{1}{2}H} \\ - Cl & 0^{-\frac{1}{2}H} \\ \end{array} \right. \\ + H^{\frac{1}{2}-0}Cl + H^{\frac{1}{2}-0-\frac{1}{2}H} \\ - Cl & 0^{-\frac{1}{2}H} \\$$

Since chlorine is negative in hydrogen chloride, it follows that the chlorine atoms in silicon tetrachloride are also negative. Of course, the hypothetical ortho-silicic acid loses water yielding meta-silicic acid, which in turn gives the anhydride, silicon dioxide—

In silicon hydride and silicon tetrachloride, hydrogen and chlorine respectively function negatively. Compounds and reactions will now be considered in which chlorine functions positively

B. Positive Chlorine.

It has been noted (p. 15) that positive chlorine ions may arise during the electrolysis of hydrochloric acid through anodic oxidation of negative chlorine ions and precede the formation of molecular chlorine. Apart from the phenomena of electrolysis, positive chlorine ions may exist and as such function in the production of hypochlorous acid through the interaction (reversible) of chlorine and water according to the equation,

which is represented ionically and in terms of electronic formulas as follows:—

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The chloime atom in hypochlorous acid is positive and the instability of hypochlorous acid and its oridizing properties may be attributed to the tendency of positive chlorine to revert to the more stable negative chlorine. It has been shown that negative rhydrogen acts as a reducing agent through the loss of electrons

and thereby reverts to positive hydrogen: $\tilde{H} - 2 \bigoplus \rightarrow \tilde{H}$ On the other hand, positive chlorine acts as an oxidizing agent through the acquisition of electrons and thereby reverts to negative chlorine.—

The decomposition (accelerated by the action of light) of dilute aqueous solutions of hypochlorous acid, yielding fice oxygen and hydrochloric acid, according to the equation, $2 \text{HOCl} \rightarrow 2 \text{HCl} + O_{\text{b}}$ may be interpreted ionically and electronically as follows:—

(1)
$$\Pi \stackrel{+}{\longrightarrow} \stackrel{-}{\longrightarrow} C \stackrel{+}{\longrightarrow} \stackrel{+}{\longrightarrow} \stackrel{+}{\longrightarrow} \stackrel{-}{\longrightarrow} \stackrel{+}{\longrightarrow} C \stackrel{+}{\longrightarrow}$$

Equation (1) indicates complete ionic dissociation of hypochlorous acid. In (2) the oxygen ion through the loss of two electrons is oxidized to the atomic or electrically neutral state

O, or O This oxidation is effected by the positive chlorine ion, Cl, which through the acquisition of two elections becomes a negative chlorine ion as noted in (3) In (4) the resultant negative chlorine ion unites with the positive hydrogen ion from (1), to give hydrogen chloride. This detailed interpretation may be abbreviated to conform to the scheme of an intramolecular oxidation-reduction reaction:—

The first electromet, $H^+ - O^- + Cl$, is converted to the second electromer $H^- - O^+ - Cl$, through the reduction of Cl to Cl and the concomitant oxidation of O to O. The second electromer, $H^+ - O^+ - Cl$, yields $H^+ - Cl$ and molecular oxygen, $O^+ - O$. Another electronic formula for oxygen is possible, namely, $O^+ - O$, but it is irrelevant to the explanation which formula is assigned to molecular oxygen.

In presenting these interpretations, it should be remembered that it is no more possible to picture the absolute mechanism of a chemical change than it is to present a final explanation in science. Each of the preceding interpretations of the decomposition of hypochlorous acid involves one and the same fundamental principle, namely, the transition of positive chlorine to negative chlorine through the loss of electrons. Furthermore, this transition serves to explain the oxidizing action of hypochlorous acid. In some reactions (to be described later) hypochlorous acid yields through dissociation, negative hydroxyl and positive chlorine ions —

Oxidation is effected through the reduction of \vec{C} l to \vec{C} l as noted above. In other reactions, hypochlorous acid undergoes intamolecular oxidation-reduction and yields neutral or atomic (the so-called "nascent") oxygen, \vec{O} , which in turn effects oxidation through its reduction to \vec{O} . In other words, positive chlorine or neutral oxygen atoms effect oxidation through transitions (1) and (2) respectively:—

In terms of the electronic conception of positive and negative

valences, transition (I) is fundamental to the decomposition of

Since the acquisition of a given number of negative electrons is electrically equivalent to the loss of the same number of equivalent positive charges, transitions (1) and (2), respectively, may be indicated thus:—

Apropos of the electronic formula and properties of hypochlorous acid, the existence of positive chloring rons is substantiated by the apparently anomalous fact that on electrolysis concentrated solutions of hydrochloric acid yield equal volumes of hydrogen and chloring at cathode and anode respectively, but, on the other hand. very dilute solutions of the acid yield hydrogen at the cathode but no free chloring at the anode. Instead an equivalent amount of arveen is evolved. Ostwald 20 states that "this is due to the fact that the water is decomposed by the chloring with the formation of hydrogen chloride and oxygen according to the equation 2H.O + 2Cl. = 4HCl + O. This process, it is true, takes place with measurable velocity only in the light: we may however, assume here as in similar cases. that the process takes place without light, only very slowly," Ostwald's explanation should be modified and extended because he has not taken into account the fact that the liberation of oxygen from a solution of chlorine in water is due, most likely, to the intermediately formed hypochlorous acid according to the generally accepted equations :-

$${}_2Cl_2 + 2HOH \Rightarrow 2HCl + 2HOCl;$$

 ${}_2HOCl \Rightarrow 2HCl + 0_m$

In view of the electronic interpretation of the liberation of oxygen from hypochlorous acid, it follows that the evolution of oxygen at the anode during electrolysis of very dilute solutions of hydrochloric may be due to the following changes. Negative chlorine ions are oxidized at the anode to positive ions:

$$CI \rightarrow CI + 2 \bigcirc, \text{ or } CI + 2 \bigcirc \rightarrow CI$$

The dissociation of water, though extremely slight, yields hydrogen and hydroxyl ions which latter migrate to the anode and there encounter a much greater concentration of positive chlorine ions. The production of hypochlorous acid through the union of the negative hydroxyl and the positive chlorine ions, and its decomposition, as previously explained, fully accounts for the liberation of oxygen. Or, in equivalent terms, this change may readily take place through the oxidation of the hydroxyl ions by positive chlorine ions at the anode. Hus :—

or summarized briefly, * $_2HO + _2CI \rightarrow _2H + _2CI + O.$

It is evident that this oxidation would proceed only in very dute solutions of hydrochloric acid, i.e., in the presence of a very low concentration of hydrogen ions. Concentrated solutions of hydrochloric acid yield no oxygen on electrolysis because the high hydrogen ion concentration obliterates the

hydroxyl ion concentration $(H + O\bar{H} \rightarrow H_2O)$. Hence in the absence of hydroxyl ions, the positive chlorine ions combine with the negative chlorine ions yielding molecular chlorine. In this way, the electronic conception of valence affords an interpretation of the apparently anomalous results obtained in the electrolysis of dilute and of concentrated solutions of hydrogen chloride.

C. Chloramines.

Not only hypochlorites but other compounds contain positive chlorine. Selivanow 22 noted the fact that the chloramines,

* In simpler terms this change may be regarded, in a final analysis, as the action of positive chlorine ions upon negative oxygen ions —

In this connection it is noteworthy that Nernst 21 regards the action of chlorine upon oxygen ions, molecular oxygen being liberated, as analogous to the action of chlorine upon bromine ions, molecular bromine being liberated. In terms of electronic formulæ this latter change is reduced to the equation—

$$(Cl_2 = Cl.Cl) + 2Br \rightarrow (Br.Br = Br_2) + 2Cl.$$

RNHCl and R₄NCl, on hydrolysis and interaction with hydrogen iorilde suffered replacement of their chlorine atoms, not by hydroxyl, but by hydrogen, and that two equivalents of iodine were liberated for each equivalent of replaced chlorine, thus

Selivanow attributed this remarkable reaction to the fact that the chlorine atom in chloramine existed as "hypochlorous chlorine". Accordingly, the above reaction is perfectly analogous to the action of hypochlorous acid upon hydrogen iodide —

In terms of electronic formulæ, hypochlorous chlorine is positive chlorine,* and, therefore, the chloramine should yield positive chlorine on dissociation —

$$R_aNCI = R_aN \xrightarrow{-+} CI \Rightarrow R_aN + CI$$

Hence in the above reaction, $\overset{\circ}{\Gamma}$ 1 oxidizes the negative iodine ion of the hydriodic acid to $\overset{\circ}{\Gamma}$ which in turn unites with $\overset{\circ}{\Gamma}$ to form molecular iodine. An analysis and summary of these changes is given in the following scheme: -

Accordingly, the analogous reactions (1), and (2),

(1)
$$R_2N \cdot Cl + 2H \cdot I \rightarrow R_2N \cdot H + H \cdot Cl + I_2$$

(2) $HO \cdot Cl + 2H \cdot I \rightarrow H \cdot OH + H \cdot Cl + I_n$

are represented respectively in abbreviated electronic formulæ as follows:—

(1)
$$\overrightarrow{R_2N}$$
, \overrightarrow{C} + $2\overrightarrow{H}$, \overrightarrow{I} \rightarrow $\overrightarrow{R_2N}$, \overrightarrow{H} + \overrightarrow{H} , \overrightarrow{C} + \overrightarrow{I} , \overrightarrow{I}
(2) \overrightarrow{HO} , \overrightarrow{C} + \overrightarrow{O} , \overrightarrow{H} + \overrightarrow{H} , \overrightarrow{C} + \overrightarrow{I} , \overrightarrow{I}

^{*} The autoseptic action of Dakin's solution and his chloramines is undoubtedly due to positive chlorine.

Each of these reactions involves one and the same change, namely .--

The existence of positive chlorine in the chloramines is further attested by a reaction for their preparation, namely, the action of hypochlorous acid upon amines. These reactions have been shown to be reversible:—

Furthermore, molecular chlorine is liberated when the chloramines interact with hydrogen chloride :--

Other remarkable properties of the chloramines which have a direct bearing upon the constitution and electronic formula of benzene will be considered later

D. Positive Bromine, Iodine, and Cyanogen.

Halogens other than chlorine manifest the property of electronic amphotesism. This is shown by a number of hydrolytic reactions. Thus tetrabromomethane and tetraiodomethane yield bromoform and lodoform respectively:—

$$Br_0C$$
, Br + H , OH \Rightarrow Br_1C , H + HO Br
 LC , I + H , OH \Rightarrow LC , H + HO , I .

Hypobiomous and hypoiodous acids respectively embody positive bromine and positive iodine.

Iodine monochloride yields hydrogen chloride and hypoiodous acid in which chlorine and iodine are respectively negative and positive:—

Nef's ²³ observations that (1) chloiocyanogen yields hydiogen chloride and cyanic acid, while (2) iodocyanogen gives hypoiodous and prussic acid, show that the cyanogen iadical, similarly to the halogen atoms, may function either positively or negatively.—

(1)
$$\overrightarrow{Cl}$$
 . \overrightarrow{CN} + \overrightarrow{H} . \overrightarrow{OH} \rightarrow \overrightarrow{H} . \overrightarrow{Cl} + \overrightarrow{HO} . \overrightarrow{CN}

$$(2]$$
 $\stackrel{+}{\text{I}}$ $\stackrel{-}{\text{CN}}$ $\stackrel{+}{\text{H}}$ $\stackrel{-}{\text{OH}}$ \rightarrow $\stackrel{-}{\text{HO}}$ $\stackrel{+}{\text{I}}$ $\stackrel{+}{\text{H}}$ $\stackrel{-}{\text{CN}}$.

14 THE ELECTRONIC CONCEPTION OF VALENCE

Thus, there are numerous instances amongst compounds classed as non-electrolytes in which polarity of atoms and radicals is definitely manifested. These examples are readily interpreted and correlated in terms of the electronic conception of positive and negative valences. The reactions just cited are but a few of the noteworthy examples of electronic amphoterism. Ordinary equations and customary formulæ do not reveal the significance of these relationships.

CHAPTER V

THE NASCENT STATE.

THE so-called "nascent state" is mitinately related to, and may be interpreted in terms of electronic amphoterism. Nascent action, according to common usage, is a term²⁴ for "all those phenomena in which a substance at the moment of its liberation from compounds performs reactions it is incapable of in its ordinary condition". Thus, to cite a simple and well-known case hydrogen has no action on silver chloride suspended in a "quid through which it is bubbled, hydrogen evolved within the liquid, electrolytically or by action of metals, produces metallic silver, hydrochloric acid being formed at the same time.

Concerning the various interpretations which have been proposed for such reactions, Alexander Smith 2 writes as follows. The term nascent hydrogen is used in different senses in a very confusing way. (i) It may mean nascent, literally, that is, newly born or liberated. (2) It is used also to mean different-from-ordinary, or, in fact, an allotropic form of hydrogen. (3) It is often limited to mean one particular allotrope, namely, atomic hydrogen. (4) It is used by Haber and others . . . to mean hydrogen activated by contact with a metal. (5) Finally, its activity is explained as being due to the larger amount of free energy contained in zinc plus acid plus reducing agent, as compared with the free hydrogen plus reducing agent. The last is identical with the explanation of the activity of exidizing agents. The word nascent is, of course, a misnomer, excepting in connection with (1) 2 metals.

A more comprehensive and conciliatory view of the nascent state and of nascent action may be developed in terms of the electronic conceptions of oxidation and reduction. Ordinarly, hydrogen functions positively while chlorine functions negatively, but in certain compounds, or under special conditions, previously described, hydrogen may function negatively and chlorine positively. Furthermore, negative hydrogen has been shown to act as a reducing agent since it naturally tends to revert to its more stable state, positive hydrogen, through the liberation of electrons (1):—

$$\stackrel{-}{\text{H}} \ \Rightarrow \ \stackrel{+}{\text{H}} \ + \ 2 \ \bigcirc .$$

On the other hand, positive chlorine acts as an oxidizing agent, since it naturally tends to become negative by acquisition of electrons (2).—

A closer analysis shows that an electrically neutral state, i.e., the atomic state, is a condition to be considered in the above transitions (1) and (2). This is shown in the following schemes (3) and (4) respectively —

(3)
$$H \rightarrow H + \bigcirc$$
, $H \rightarrow \overset{+}{H} + \bigcirc$.
(4) $\overset{+}{Cl} + \bigcirc \rightarrow \overset{-}{Cl}$; $Cl + \bigcirc \rightarrow \overset{-}{Cl}$.

Now, if the atomic state and the mascent state are to be regarded as synonymous, then they may be defined, from the electronic standpoint, as an unstable condition which manifests not only an adaptability but also a tendency either to gain or to lose electrons and thereby attain a more stable condition. This tendency, however, can not be limited to the atomic state of an element, since not only atomic or neutral hydrogen, but also negative hydrogen tends to lose electrons. A survey of equations (1) and (3) indicates that a negative hydrogen atom may be more potent as a reducing factor than a neutral hydrogen atom since the former yields two electrons and the latter only one in the transition to the positive state. The activity of negative hydrogen as a reducing agent has been illustrated by the action of sodium

hydride (Na. H) upon carbon dioxide. The product of the reduction is sodium formate. It may also be concluded that either negative or neutral hydrogen atoms produced at the cathode may function as the active reducing agent in the socalled electrolytic reductions

On the other hand, positive chlorine is a more potent oxidizing agent than is atomic or neutral chlorine since the former unit may combine with two electrons while the latter unites with only one in the transition to negative chlorine, as indicated in equations (2) and (4) above. It has just been shown, in the preceding chapter, that the formation of positive chlorine ions at the anode would account for the liberation of oxygen during the electrolysis of very dilute solutions of hydrochloric acid (low hydrogen ion concentration) according to the abbreviated equation —

Therefore, from these briefly developed points of view, the masent state may be defined more comprehensively as an unstable most of a substance which monifasts both an adaptability and a tendency to lose electrons, or to gain electrons, and thereby revol to a more stable condition. If the substance (ion, atom, or compound) loses electrons it acts as a reducing agent. If it combines with electrons, it acts as an axidizing agent. As a matter of fact, practically all actions classed as "nascent" are of an oxidation or a reduction type.

This conception of the nascent state serves as the basis for an interpretation of the existence and the properties of "free radicals" to be fully considered in a later chapter. It will be shown that free radicals may be defined as electrically neutral atoms or malecules which are capable of developing either positive of negative valences depending upon certain conditions which affect the loss or gain of electrons

CHAPTER VI.

IONIC AMPHOTERISM

ELEMENTS which gain or lose electrons and thereby function in different ways are termed amphoteric. Compounds are designated as amphoteric when, through different modes of ionization, they function either as acids or as bases. Accordingly, electronic amphoter ism relates to elements while ionic amphoterism relates to compound. It is essential to subsequent developments to consider briefly the phenomena of ionic amphotenism from the point of view of the electronic formulæ of amphoteric compounds.

While hydrogen may function negatively and oxygen positively, it will be found that in the electronic formulae of mocompounds the hydrogen atoms manifest one positive valence while the oxygen atoms display two negative valences. The simplest illustration is that of water, $H \stackrel{+}{\longrightarrow} U \stackrel{-}{\longrightarrow} U \stackrel{+}{\longrightarrow} U$

Ionic amphoterism ²⁶ is manifested by both acids and bases (metal hydroxides). Stieglitz ²⁷ maintains that "pronounced amphoterism is shown by a large number of metal hydroxides, it is, perhaps the rule rather than the exception". It should be added that many acids also show pronounced ionic amphoterism as is evident from the distinctive types and products of interaction. For example, hypochlorous, nitric, and sulphuric acids yield not only (1) positive hydrogen ions but also (2) negative hydroxyl ions. From this point of view, the electronic formulæ of these acids and their ionization, either (f) as an acid, or (2) as a base, respectively, are indicated in the following schemes:—

A general electronic formula may now be employed for each of these acids, namely, $H \xrightarrow{+} O \xrightarrow{-} X$, m which $\overset{+}{X} = C_{l_1}^{+} NO_{2l_2}^{+}$ or $SO_{a}H$.

In neutralization reactions HOX, dissociating as an acid, interacts with bases, say NaOH, according to the ionic scheme for double decomposition reactions—

On the other hand, when hypochlorous acid acts as a chlorinating agent, or when nitric acid acts as a nitrating agent, or when sulphuric acid functions as a sulphonatung agent, the acid dissociate as hases The chlorination, nitration, or sulphonation of benzene likewise would conform to the ionic scheme for a double decomposition reaction —

$$H^{+}_{-0}^{-+}X \Rightarrow H_{0} + X$$
 $C_{0}^{-}H_{0}, H^{+} \Rightarrow H^{+}_{-0} + C_{0}^{-}H_{0}$
 $\downarrow \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \qquad \qquad \qquad$

The preceding schemes illustrating (1) acid and (2) basic dissociation of HOX are summarized in the following reactions:—

(t)
$$\overset{+}{H}.OX + \overset{+}{N_{A}}.OH \rightarrow \overset{+}{H}.OH + \overset{+}{N_{A}}.OX$$
.
(2) $\overset{-}{H}O.X + \overset{+}{C_{B}}H_{B}.H \rightarrow \overset{+}{H}.OH + \overset{-}{C_{B}}H_{B}.X$

It should be noted that in direct substitution reactions, i.e.,

when X replaces H as in equation (2), $\overset{+}{X}$ usually replaces $\overset{+}{H}$. Such a change involves neither oxidation nor reduction. The substituent, however, may, subsequently, under certain conditions, function negatively. In such instances (to be presented later) the compound C_tH_s , X manifests the phenomena of electronic tautomerism $(\overset{+}{C}_0H_s,\overset{+}{X})$ $\rightleftharpoons \overset{+}{C}_0H_s,\overset{-}{X})$.

The oxidizing action of certain oxygen acids is due undoubtedly to their capacity to ionize as bases. The oxidizing action of hypotholrous acid has been explained in terms of its basic ionization, $HO : Cl \Rightarrow HO + Cl$, and the tendency of Cl to severt to Cl, thus .—

The oxidizing action of nitric acid may be attributed to basic dissocation. For example, the oxidation of a ferrous salt is usually represented by the molecular equation:—

$$2FeSO_4 + H_9SO_4 + 2HNO_3 \rightarrow Fe_2(SO_1)_3 + 2IL_9O + 2NO_3$$

If the nitric acid ionizes as a base, $HO \cdot NO_2 \Rightarrow HO + NO_3$, then the above equation may be written ionically as follows —

$$^{++}$$
 $^{--}$ $^{+}$ $^{--}$ $^{++++}$ $^{--}$ $^{--}$ $^{++++}$ $^{--}$ $^$

Eliminating the ions common to each side of the above reaction as well as the H and OH ions which form water, the following simple equation remains:—

The positive ion, NO₂ (resulting from the basic dissociation of nitric acid) is discharged by an electron from the ferrous ion, $\stackrel{+}{\text{Fe}}$ + . Accordingly, $\stackrel{+}{\text{NO}}_{\text{q}}$ is reduced to neutral or molecular

NO₂ which passes off as a brown gas. Concomitantly, Fe is oxidized to Fe⁺⁺⁺. If sulphuric acid were not used in the above reactions, its part in supplying positive hydrogen ions would be played by the acidic ionization of the nitric acid.

As to sulphuric acid, when copper interacts with hot concentrated sulphuric acid according to the equation $Cu + 2H_2SO_1$ $- CuSO_1 + 2H_2O + SO_2$; one molecule of the sulphuric acid undergoes (1) acidic ionization, the other undergoes (2) basic ionization and (3) the metal copper becomes ionic through the loss of two electrons —

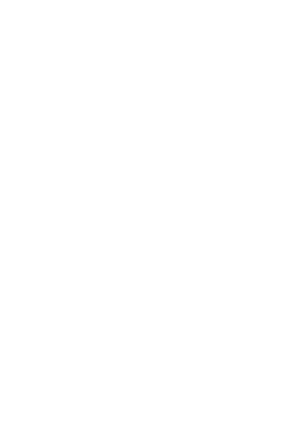
The positive ion, SO₂, is reduced to the neutral state and liberated as sulphur dioxide gas. Concomitantly, metallic copper is oxidized to the ionic state, Cu ++. The summation of (11, (2), and (3) gives the complete equation for the reaction.

Many other reactions are readily interpreted in terms of the ionic amphoteism of acids Subsequent applications of the conception of positive and negative valences will involve the principles of electronic and ionic amphoteism, and the nascent state as outlined in this and the preceding chapter



PART II.

THE ELECTRONIC FORMULA OF BENZENE: SUBSTITUTION IN THE BENZENE NUCLEUS.



CHAPTER VII.

THE CONSTITUTION OF BENZENE.

A, The Benzene Theory.

In 1865 Kekulé ** proposed his well-known formula for benzene, involving the assumptions that the six carbon atoms in benzene form a closed chair or nucleus, that the indicate is premartical, and that each carbon atom is directly united to the same only one atom of hydrogen. These assumptions are embodied in the partial structural formula.—

which, however, does not take account of the fourth unit of valency of the carbon atoms in the nucleus. The disposition of the fourth valence has been the occasion of extended discussion and speculation.

Since 1865 a variety of benzene formulae have been proposed, foremost of which are the following: the unsymmetrical formula of Str James Dewar (1865), the diagonal formula of A Claus ³⁰ (1867); the prismatic formula of A Ladenburg ³¹ (1869), the "centric formula" of H E. 'Armstrong ³² (1887), and Baeyei ³² (1888). In addition to the foregoing plane formula, a number of stereochemical configurations of the benzene complex are noteworthy: the octahedral formula of Julius Thomsen, "(1886); W. Vaubel's ³⁰ configuration (1894); the Sachse ³⁶ model (1888); the dynamic formulae of J. N. Colhe ²⁷ (1897), and the device of B. Konig ³⁶ (1995)

It is not the purpose of this monograph to discuss the characteristic features of the foregoing benzene theories. Their

respective menits or demerits can be determined by a careful study of the references cited, which will reveal the fact that none of the proposed formulae is accepted as complete H. E. Armstong ⁵⁰ summarizes the stuation as follows: "The determination of the 'structure' of this hydrocaubon [benzene] has given rise to a large amount of paper warfare. Two tendencies may be said to have been brought out in the course of the discussion: on the one hand, the desire to arrive at a determination of the actual structure, on the other, the desire to devise formulae which shall be faithful to the functional behaviour and broadly indicative of the structural relationship of the constituent elements. The latter is perhaps the tendency which is now in the accendant: we are beginning to realize particularly in the case of carbon compounds that formulae are primarily expressive of chavarous—behavi

Notwithstanding these developments, we are still confronted with the fact that none of the proposed formulæ for benzene has lent itself to a uniform systematic explanation of substitution in the benzene nucleus and the many anomalous reactions of benzene and its derivatives. The foremost authority on the problem of substitution in the benzene nucleus, A. F. Holleman, and has recently written as follows. "Notwithstanding the fact that the problem of substitution in the benzene nucleus has been studied intensively enough of late, there still remains a fundamental question which has not yet been solved, it is the question of knowing the reason why such or such group directs a new substituent chiefly to the para-ortho positions or chiefly to the mean position."

The great variety of benzene formulæ and their marked deniciencies in interpreting and con elating the behaviour of benzene and its derivatives, especially the substitution reactions, has led to the generally accepted conclusion that the constitution of the benzens nucleus presents a remarkable case which must be dealt within the formulation of any complete theory of valence. Hence, a critical test of the utility of the electronic conception of positive and negative valences may be made by applying the principles developed in the preceding chapters to the constitution of the benzene nucleus. An electronic formula for benzene is thereby derived. No claim whatever is made that this electronic formula is the final solution to the manifold problem of the constitution of benzene, but it will be shown that the electionic formula of benzene and the underlying principles of the electionic conception of valence afford a means of interpreting and correlating many litherto unexplained chemical, physico-chemical, and stereochemical problems presented by benzene and its derivatives,

B. The Electronic Formula of Benzene.

In the benzene formulæ proposed by Kekulé, Claus, Armstrong, and Baeyer, respectively, two of the units of valency of each of the six carbon atoms function in the formation of a closed homocyclic ring or "benzene nucleus". The third unit of valency engages a hydrogen atom. The disposition of the fourth unit of valency led Kekulé to propose the existence of three "olefin" or double bonds in the ring. It is conceded that these are not of the same nature as the double bonds in the hydrocarbons of the olefin series. In the Claus and Ladenburg, formulæ, the fourth unit of valency, or bond, is represented as directly united to three other carbon atoms. In the Aunstrong and Baever "centic" formulæ the fourth unit of valency is directed towards a centre This signifies that the six carbon atoms of the ring' have a general attraction for one another but they are not directly united in the usual way by the fourth unit of valency. To what configuration, then, is the electronic conception of valence to be applied in order to derive an electronic formula for benzene? The configurations just noted are plane and static a complehensive benzene configuration must be stereochemical and dynamic.

In a discussion of the ments of the various space formulæ of benzene, Stewart " states that "the objections brought against the older types of space formulæ show that any advance in this branch of the subject must follow the lead given by Kekulé when he adopted the idea of a system in wibiation as the best representation of the benzene molecule." At the present day, the idea that benzene is one particular substance which can be represented at all times and under all conditions by the same rigid formula finds very few supporters among, those who have studied the question thoughly. It is becoming generally recognized that the benzene molecule is in a state of continual vibration, and that the only satisfactory space formula will be now which represents all the other formula as phases of its own

motions, and which may even suggest the possibility of new phases as yet uniccognized. The main outlines of such a formula have been indicated by Collie, i* and it seems probable that any space formulae of benzene which may be proposed in the future will agree with his in essentials."

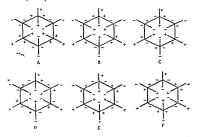
Collie's well-known space formula of benzene comprises a system of six caibon atoms (tetrahedra) each united to a hydrogen atom. Movement in this system can take place in two ways: (1) Movement of each tetrahedron about its centre, (2) movement of each tetrahedron about the centre of gravity of the whole system. The plane projections of the symmetrical configurations rendered possible are represented by Collie as follows:—

As stated by Stewart, "" "this space formula for bertzene, herefore, is in complete accord both with the Kekulé and the centric formula; showing that they are mutually convertible into one another. It also shows how the supposed double linkages of the Kekulé formula shift between the carbon atoms, rendering the existence of two o-toluidmes impossible. But it differs from both in that in two out of five configurations there are two distinct sets of hydrogen atoms." . The Collie formula are "in perfect accord with the formula of Ladenburg, Claus, or Dewar, or that of Baly, Edwards, and Stewart. This adaptability is not possessed in any degree by previous space formulae, and it is this which makes the Collie formula superior to the others."

If this pre-eminence be granted to the Collie space formula, then it is the formula to which the electronic conception of valence should be applied in order to derive the most comprehensive electronic formula for benzene. In making this application it will be sufficient, for the present, to consider in detail the constituent aloms of only one of the five interconvertable phases, of the plane projections of Collie's space formula, namely, the centric formula. The derivation of an electronic formula from the centric formula will now be considered.

The quadrivalent carbon atom may function electronically in five different ways ---

If benzene nuclei (centric formule) are composed of these several types of carbon atoms, each nucleus consisting of three pairs of the combined types I and V., II, and IV., and III, and III, symmetrically co-ordinated, then six and only six centric electronic formulae, or electromers, are possible. They are as follows (A—P)—



It has been noted that there are five plane projection formulas of the Collie space formula to benzene of which the centric formula is one. Now since there are six centre electronic formulae for benzene, (A—F), it follows that there are six times as many electronic formulae for benzene as there are plane formulae in the Collie system. If we disregard the double bonds of the Kekulé formulae and the centric bonds of the centric

formulæ, an abbreviated electronic formula for benzene is represented thus:-

This formula is typical of each of the possible electronic formula or electromers of benzene. While it has been derived in an a piont manner, so to speak, thue are, nevertheless, pronounced reasons for its proposal. These reasons are embodied in the following distinctive characteristics.

- (1) If the hydrogen atoms in positions 1, 3, and 5 are negative, those in positions 2, 4, and 6 are positive. Or, since the electronic formula is perfectly symmetrical, if the hydrogen atoms in positions 1, 3, and 5 function positively, then those in positions 2, 4, and 6 function negatively.
- (a) The Kekulé and Centric formulæ of benzene fail to show any structural basis for the relationships existing between orthoand para-positions in contradistinction to the meta-positions. The electronic formula shows at once that if a given hydrogen atom or substituent is negative, then those hydrogen atoms or substituents ortho and para to it are each positive, while those meta to it are each negative. Or, if a given hydrogen atom or substituent is positive, then those atoms or substituents ortho and para to it are each negative, while those meta to it are each positive.
- (3) From the foregoing, it is evident that the electronic formula for benzene presents, per se, the basis for a definite substitution rule, namely, when substituents are of the same sign or polarity they will occupy positions which are meta to each other: if two substituents are of opposite sign or polarity they will occupy positions either with or para to each other.
- (4) Collie's space formulæ for benzene, the plane projections of which have been noted, show, in two of the five phases, the existence of two sets of hydrogen atoms, namely, the I, 3, 5- and the 2, 4,6-set Concordantly, there exists in every electronic formula.

Therefore, the entering positive halogen atom, in substituting a positive hydrogen atom, must likewise occupy a position meta to the positive carboxyl radical. This reaction and the electronic formula of benzene are thus correlated. Furthermore, the dissociation of the halogen molecule into positive and negative parts, as indicated in the above equation, is a confirmation of the idea suggested at an eather date by W. A. Noves 11 that "if we suppose what seems not to be improbable, that all reactions involving the decomposition of molecules are preceded by an ionization of the parts of the molecules, it would follow that elementary molecules as well may ionize into positive and negative parts".

C. Transference of Radicals from the Side-Chain to the Nucleus.

Various isometic changes, stereochemical rearrangement reactions, may be explained in terms of the electronic conception of valence and the electronic formula of benzene The remarkable rearrangements characteristic of the substituted nitrogen halides investigated by Chattaway and Oiton 15 are of particular interest. For example, acetanilide on treatment with hypochlorous acid yields phenylacetylnitrogenchloride which is readily transformed into hara-chloroacetanilide. This on treatment with hypochlorous acid gives para-chlorophenylacetylnitrogenchloride which is transformed into 2: 4-dichloroacetantide. Proceeding thus, the final product is 2:4:6-trichloroacetanilide which will not undergo rearrangement. These successive changes are summanzed in the following scheme in which R = formyl, acetyl, or benzovl. X = Cl or Br .-

Why does the entering halogen atom invariably migrate to

a position in the nucleus para or ortho to the amino group, provided these positions are unoccupied? Anılıne as a derivative of ammonia, H.———II, may be represented by the abbrevi

ated electronic formula, $C_4H_3^+--NH_p$. The radical NH_2 is negative and accordingly must occupy the position of a negative hydrogen atom of the benzene nucleus. The halogen atom in hypochlorous or hypobromous acid $(X=Cl\ or\ Br)$, has been shown to function positively and its interaction with an anilide to conform to the reaction—

The following more comprehensive electronic formula for phenylacylnitrogenhalide follows ---

Therefore, in the process of transformation into a less labile isomenicle, it is self evident that the posttree halogen atom can exchange positions only with a positive hydrogen atom of the benzene nucleus; i.e., in passing from the nitrogen atom to the nucleus it must assume either the pair or ortho positions with respect to the attachment of the NH_2- or RNH- radical which functions negativelp. When the para and both ortho positions are occupied, as in 2,4,6 throbromoscyluttogenbromde, one finds that rearrangement is theoretically impossible from the point of view of the electronic formula and it cannot be effected experimentally. This two-fold conclusion constitutes a proof of the identity of polarity of the positions 2, 4, and 6, of the benzene nucleus, each of which is occupied by a positive chlorine atom in the trichloroot itibromo-phenylacylnitrogenhalide. Furthermore, the posi-

tive chlorine atoms of the nucleus resist hydrolysis and in this respect resemble the positive chlorine atom in metachloronitro-benzene which does not undergo hydrolysis. On the other hand, recall that the chlorine atoms are negative in ortho and para chloronitrobenzenes which compounds are completely hydrolyzed yielding respectively ortho and para introbeneols

Afiline and phenol, on bromination, are quickly and quantitatively converted into 2, 4, 6- tribromonilline and 2, 4, 6- tribromophenol, i.e., the bromine atoms occupy positions in the nucleus ortho and para to the amino and hydroxyl radicals in the respective compounds. These reactions are identical in their nature to the rearrangements which characterize the substituted nitrogen halides. The explanation of these changes in terms of electronic formulæ are also identical in principle and method. This follows from the exact similarity existing between the electronic formulæ of annulæ and planol :—

The negative amino indical of aniline and the negative hydroxyl radical of phenol each contains a positive hydrogen atom which is replaced by a positive bromme atom on interaction with free bromine or hypobromous acid. On rearrangement, the positive bromine atom can assume only those positions in the nucleus which are occupied by positive hydrogen atoms these are the positions without and para to the negative animo and the negative hydroxyl radicals. Consequently, these substitutions and teanrangements, following in rapid succession, conform to the scheme indicated for the substituted introgen haldes, the final products are 2, 4, 6- tribromo-abenol.—

Many other rearrangements involving transference of radicals from the side-cham to the nucleus, i.e., from introgen to nuclear carbon, conform to the preceding interpretation. The following examples are noteworthy: Hofmann and Martins** found that the hydrechloride of aromatic methylamino compounds at 250°-350° suffered transference of the methyl group from the nutrogen to the 67tho and para carbon atoms of the nucleus. Thus, monomethylamiline yuelds ortho-toluidine and para-toluidine, and repetition of the process results finally in the formation of 2.4.6. cumlidine —

The nitrosamine of monomethylanılıne, according to Fischer and Hepp of yıelds para-nitrosomethylanılıne; and, in similar fashion, phenylnitranılıne, according to Bambeige, iii changes to ortho- and para-nitraniline. The well-known benzidine and semidine conversions, and a variety of rearrangements to ortho and para positions encountered in the synthesis of many dyestinfis conform to, and therefore, confirm the electronic formulae oblemzene and tis substitution products. It is electronic formulae only which indicate the way in which these rearrangements take place: ordinary structural formulae do not lend themselves to an interpretation of these apparently anomalous reactions.

D. Positive and Negative Carboxyl Radicals.

Some carboxyl radicals of certain derivatives of benzene readily suffer decomposition with the elimination of carbon dioxide:—

Other carboxyl radicals, under the same conditions, are quite stable. May not this difference in stability be due to a difference in the polarity of the valence which binds the carboxyl radical to the benzene nucleus. Facts will now be considered which show that the polarity of the carboxyl radical has a definite influence upon the property of the compounds, or more particularly upon the stability of the carboxyl radical itself. A compound containing two carboxyl radicals in positions ortho to each other is found in pithalic acid. According to the electronic formula for benzene one of the curboxyl radicals of the acid should be positive while the other is nearly as indicated in the formula.

Is there any experimental evidence to justify this difference in the attachment of the carboxyl groups to the nucleus? Baeyer "has shown that the $\Delta^{3.5}$ $\Delta^{-1.5}$, and $\Delta^{3.6}$ dihydrophthalic acids yield benzoic acid on oxidation. He attributes this to a shock (Ei schutterung) to the a-carbon atom which causes it to lose carbon dioxide, this effect being associated with the change from ethylene to centric linkages, thus —

Bitill, "o in a critical examination of Baeyer's "Eischutterung Theory," asks why only one molecule instead of two molecules of carbon doxide is semoved from the acids in question, and has sought to explain the chemical behaviour of the dihydro acids on the simple basis of their difference of stability, which view has the advantage of being independent of any structural hypothesis. But Buhl, according to Cohen, "a falls to paceive that

by avoiding any reference to structure as affecting stability, he is begging the question." It must then be concluded that up to the present time there is no satisfactory structural explanation as to why only one molecule of carbon dioxide instead of two is removed from the several dihydrophthalic acids investigated by Baever

The electronic formula of benzene requires that substituents occupying ortho positions to each other be of opposite polarity. Consequently in the phthalic acids one carboxyl radical functions positively and the other negatively as previously indicated. If reference be made to the electronic formula of formic and carbonic acids, it is evident that the carboxyl radical in formic acid is negative (H—COOH) while in carbonic acid it is positive (HO—COOH). Furthermore, in formic acid three of the valences of the carbon atom are positive and one is negative. Accordingly, formic acid on decomposition ordinarily yields carbon monoxide and water —

The free positive and the free negative valence of the carbon atom in carbon monoxide become neutral. In other words, carbon in carbon monoxide is bivalent —

On the other hand, in carbonic acid and its anhydride each of the four valences is positive. Accordingly, carbonic acid on decomposition yields water and carbon dioxide —

From the above it is evident that a carboxyl radical which is negative corresponds to the stage of oxidation represented by the carbon atom in formic acid, which does not lose carbon dioxide. On the other hand, a carboxyl radical which is

positive corresponds to the stage of oxidation represented by the carbon atom in carbonic acid, which very readily loses carbon dioxide. Therefore, the phthalic acids which embody a negative carboxyl radical and a positive carboxyl radical, lose only one molecule of carbon dioxide, a result of the decomposition of the positive carboxyl radical as indicated in the following scheme:-

This explanation may now be extended to the interpretation of many other anomalous phenomena involving the liberation of carboll dioxide from various substituted benzoic acids.

E. Elimination of Carbon Dioxide.

Cazeneuve 52 in a study of the decomposition of various hydroxybenzoic acids found that some of them readily lose carbon dioxide when heated with water or aniline, while others were quite stable. For instance, orthohydroxybenzoic acid (salicylic acid) when heated with aniline to 240° in a sealed tube gave phenol and carbon dioxide. Parahydroxybenzoic acid likewise gave carbon dioxide but the meta acid suffered no change. The instability of the ortho and the para acids and the stability of the meta acid may be interpreted in terms of the electronic formula of benzene and the polanty of the carboxyl radicals. These radicals in ortho- and para-hydroxybenzoic acids must function positively, i.e., they correspond to the state of exidation represented by carbonic acid and therefore may vield carbon dioxide Furthermore, if the hydroxyl radical in the hydroxybenzoic acids is negative, then the electronic formulae for the ortho- and para-hydroxybenzoic acids require that the carboxyl radicals be positive, while the carboxyl radical in the meta acid must be negative. The meta carboxyl radical being negative corresponds to the state of oxidation of formic acid and, therefore, does not lose carbon droxide. The conclustion of these phenomena with the conception of positive and negative valences is expressed in the following formulae for ortho-, para-, and meta-hydroxybenzoic acids —

From the above, a general conclusion may be drawn, namely, that a carbaxyl radical either ortho or para to a negative radical spatitive and, therefore, unstable, yielding carbon draxide when heated with water or antline. On the other hand, a carbaxyl radical meta to a negative hydraxyl radical as also negative and, therefore, stable, not yielding carbon draxide when heated with water or aniline.

The above conclusion receives remarkable confirmation by the recent and numerous experiments of Hemmelmayr 58 in an extensive research "Concerning the influence of the nature and position of substituents upon the stability of the carboxyl radical in the substituted benzoic acids". Cazeneuve estimated the relative stability of the carboxyl radicals by comparing temperatures at which different compounds eliminated carbon dioxide Hemmelmays's method is more comprehensive, in that he heated weighed quantities of various acids with either 50 c.c. of water or 50 cc. of aniline and estimated quantitatively the amounts of carbon dioxide liberated In this manner he experimented with thirty-nine differently substituted hydroxybenzoic acids. Their behaviour in general conformed to the rule indicated above. In other words, an examination of the formula showed that every acid which contained a carboxyl radical either ortho or para to an hydroxyl radical yielded carbon dioxide On the other hand, those acids in which the carboxyl group was neither ortho nor para, but only meta, to an hydroxyl rudical did not yield carbondioxide. Thirty-five of the thirty-time variously substituted hydroxybenzoic acids conformed to this generalization. Among hirteen mono, di-, and trihydroxybenzoic acids there was not one exception. Hence, it must be concluded that the electronic formula of benzene and the conception of positively and negatively functioning carboxyl radicals afford an interpretation of the anonalous behaviour of numerous and differently substituted hydroxybenzoic acids, some of which lose carbon droxide while others do not.

While it is thus evident that positively functioning carboxyl radicals lose carbon dioxide when heated with water or aniline, nothing has been found in the literature which would indicate or explain the mechanism of these reactions.

F. The Mechanism of the Elimination of Carbon Dioxide.

In attempting to explain the action of water or anilline it should be noted that each of these reagents which effect the elimination of carbon dioxide contains an unsaturated atom, i.e., oxygen may increase its valence from two to four while nitrogen, in aniline and other amines (or ammonia), may increase its valence from three to five. For instance, the combination of aniline with Hydrogen halide may be represented as follows—

Since Hemmelmayr has found that ammes other than amiline effect the elimination of carbon dioxide it is advisable now to limit this explanation to the unsaturated state of the nitrogen atom in ammes

In a recent paper ⁵¹ I have shown that the unsaturated natiogen atom in the amine, pyridine, lends itself to the elimination of hydrogen sulphide in a new reaction for the preparation of thiocarbanilides, according to the equation—

$$_{2RNH_{4}}$$
 + $_{CS_{2}}$ + $_{C_{3}H_{4}N}$ \rightarrow $_{(RNH)_{2}CS}$ + $_{C_{3}H_{4}N}$ $\stackrel{H}{\searrow}_{SH}$ $_{C_{3}H_{4}N}$ $\stackrel{H}{\searrow}_{SH}$ \rightarrow $_{C_{4}H_{4}N}$ + $_{H.SH.}$

In other words pyridine combines with the elements of hydrogen sulphide forming the unstable pyridinium sulphide which, decomposing, regenerates the original pyridine and eliminates hydrogen sulphide. Analogously, it may be assumed that an unstable addition compound of an amine and the asyberiose acid is formed which, adcomposing, regenerates the amine and theo areas carbon disadile.

The validity of this assumption was put to an experimental test by employing pyridine as the amine and chlorocatbonic ethylester as an analogue of an hydroxybenzoic acid. The possibility of the climination of carbon dioxide from chlorocatbonic ethylester is wainated, since all of the valences of its carbon atom are positive and so correspond to the state of oxidation of the carbon atom in carbonic acid and carbon dioxide. This follows from its interaction with water.—

It should be noted, parenthetically, that chlorocarbonic ethylester is very frequently, but incorrectly, called "chloroformic ethylester" which implies that it is a derivative of formic acid. If such were the case then its carbon atom must possess three positive valences and one negative valence in order that it correctly correspond to the state of oxidation of the carbon atom in formic acid. Its electronic formula would then be as follows:—

This "electromer" if hydrolyzed could not yield carbon dioxide directly. The products would be hypochlorous acid,

$$H^{+} O^{-} CI$$
, and ethyl formate-
$$H^{+} O^{-} C_{1} H_{0}$$

$$H^{+} O^{-} C_{1} H_{0}$$

but, as just noted, this is not the case.

When chlorocarbonic ethylester is dissolved in an anhydrous solvent (carbon tetrachloride, benzenc, or toluene) the addition of a like solution of pyridine yields at once an extremely unstable addition-compound (white solid, not yet isolated) which spontaneously decomposes, bloeating carbon dioxide and ethyl chloride. Experiments were performed in order to determine quantitatively the extent of the decomposation of various amounts of chlorocarbonic ethylester by a fixed amount of pyridine. The results of these experiments are embodied in the following table, which piesents the molecular ratios of the interacting substances, the quantities employed, the amount of carbon (hoxide evolved and the corresponding percentages of the theoretical yields of carbon dioxide based upon the ratio CO₂, CICO₂,CH₂.

Mulecular Ratios	C ₃ H ₅ N (Grams)	CICO ₂ C ₃ P ₁₆ (Grams)	CO ₂ (Grattis)	Per Cent of Theory
2 C ₅ H ₆ N · 1 CICO ₂ C ₂ H ₈	3 95	27t	0°9720	88:36
2 C ₅ H ₆ N · 2 CICO ₂ C ₃ H ₃	3 95	5'42	1°8960	86:18
2 C ₅ H ₅ N · 3 CICO ₂ C ₂ H ₈	3 95	8'13	2°73°5	82:71

The reaction mixture in the flask still contained pyridine which effected the decomposition of additional quantities of chlorocarbonic ethylester. Hence it must be concluded that the pyridine acts as a catalytic agent in effecting the elimination of caubon dioxide from the este through the intermediate formation of an unstable addition compound. The reactions involved may be represented by the following equations:—

$$C_{a}H_{b}N + CI \xrightarrow{+C^{+}} C \xrightarrow{-C^{+}} C_{b}H_{b} \rightarrow C_{a}H_{b} \xrightarrow{-C^{-}} C \xrightarrow{-C^{-}} C_{b}H_{b}$$
 $C_{a}H_{b}N + CI \xrightarrow{+C^{-}} C \xrightarrow{-C^{-}} C_{b}H_{b}$
 $C_{a}H_{b}N + CI \xrightarrow{+C^{-}} C \xrightarrow{-C^{-}} C_{b}H_{b}$
 $C_{a}H_{b}N + CI \xrightarrow{+C^{-}} C \xrightarrow{-C^{-}} C_{b}H_{b}$
 $C_{a}H_{b}N + CI \xrightarrow{+C^{-}} C \xrightarrow{-C^{-}} C_{b}H_{b}$

I have found that amines, other than pyridine, also effect the elimination of carbon dioxade from chlorocarbonic methyl, ethyl, and propyl-esters. Hemmelmayr has found that amines other than aniline effect the elimination of carbon dioxade from various hydroxybenzole acids in which the carboxyl radicals have been shown to function positively. Therefore, equations perfectly analogous to those above may represent the action of an amine of the general formula R₂N upon an hydroxybenzole acid (so any other substituted benzole acid in which the carboxyl radical is positive) as follows: (Equations showing the formation of the amine salts of the acids have been omitted as they are not essential to the puncipal changes involved).

(R = Alkyl, aryl, or H; X = OH or other substituent.)

The formation of an intermediate unstable compound of the amine and the acid depends upon an increase in the valence of the nitrogen from three to five. Similarly, the action of water in effecting the elimination of carbon dioxide would depend upon the formation of an unstable oxonium compound though an increase in the valence of the oxygen atom from two to four as indicated in the following equations, which are perfectly analogous to those above —

Thus the electronic conception of positive and negative valences not only indicates the nature of the radicals which, though the action of amines or water, lose carbon dioxide but it also affords a possible interpretation of the way in which the reactions proceed. The theoretical deductions are based upon

experimental facts and data afforded by the analogous action \$\displainter{c}\$ pyridine upon chlorocarbonic ethylester. It is quite conceivab! that the elimination of carbon dioxide from carbamino compound of the proteins in physiological processes may conform to the type of reactions just described.

G. Positive and Negative Nitro Groups: the Hydrolysi of Nitro Derivatives of Benzene.

In a compound of the general electronic formula H--X the atom or radical X is negative; but in $H \xrightarrow{+-} O \xrightarrow{-+} X$ it i positive For instance, in nitrous acid, H.NO., the mtro grou is negative, but in nitric acid, HO NO, it is positive.

It is generally conceded that nitrous acid may exist in tw tautomene forms, presumably in equilibrium -

The electronic formulæ of tautomers I and II follow :-

In formula I, the nitrogen atom possesses one negative at four positive units of valence, the algebraical sum of which three positive units. If the hydrogen atom is transferred fro the nitiogen to the oxygen atom through the opening up of oi of the double bonds between nitrogen and oxygen, the following intermediate structure may result ---



It is assumed that polarization of the free positive and free fregative unit of valence leaves the nitrogen atom in resultant tautomer II, with three positive units. Thus, it is evident that in each of the tautomers, I, and II., the nitrogen atom possesses algebraically three positive units of valence, and this marks the state of oxidation of the intogen atom in introduced and the nutries. In either tautomeric form the NO₂ or ONO radical functions negatively.

On the other hand, in nitric acid and the nitrates the nitrogen atom is in the state of oxidation represented by five positive units of valence as indicated in the formula—

Here the NO. radical functions positively.

It may now be shown that the benzene nucleus may embody both positive and negative nitro groups, and that the latter, in hydrolytic reactions, yield nitrous acid or nitrites.

Phenol, when directly nitrated, yields ortho- and para-nitrophenol Further nitration yields the two isomenc I, 2, 4- and I, 2, 6-dinttophenols. Finally, nitration of either of these products gives the symmetrical trinitro derivative, picric acid. These reactions are summarized in the following electronic equation:—

$$\begin{array}{c} \stackrel{\bullet}{\text{OH}} \\ \stackrel{+}{\text{H}} \\ \stackrel{+}{\text{H}} \\ \stackrel{-}{\text{H}} \\ \stackrel{+}{\text{H}} \end{array} + {}_{3}\stackrel{\bullet}{\text{Ho}} \cdot \overset{+}{\text{No}_{1}} \Rightarrow \begin{array}{c} \stackrel{\bullet}{\text{No}_{1}} \\ \stackrel{\bullet}{\text{Ho}} \\ \stackrel{+}{\text{Ho}} \\ \stackrel{+}{\text{Ho}} \end{array} + {}_{3}\stackrel{\bullet}{\text{Ho}} \cdot \overset{\bullet}{\text{OH}}.$$

In tuntrophenol each of the nitro groups is positive, but the electionic formula indicates that in tetranitrophenol a fourth nitro group in position 3 or 5 would be negative. Now if some nitro groups are positive and others are negative these should be some maked difference in their behaviour toward certain reagents. These conclusions are substantiated by the fact that, as stated by J. B. Cohen, so 'the nitro group in nitrobenzene is extremely stable compared with the fourth nitro group in tetra-

nitrophenol, which water will remove in the form of introus acid? These facts are in perfect agreement with the electronic formula of tetranitrophenol, the hydrolysis of which yields introus acid. $\overset{+}{H}$, $\overset{-}{NO}_{\infty}$ or $\overset{+}{H}$, $\overset{-}{NO}NO$) according to the electronic equation:—

Since direct intration with nitric acid results in the substitution of positive into groups, and since in terraintrophenol the negative nitro group may occupy either of the equivalent positions 3 or 5, with negative hydroxyl in position 1, it follows that substituents in positions 1, 3, and 5 are negative while those in positions 2, 4, and 6 are positive. This is a confirmation of the electionic formula of benziene and the substitution rule.

It has been shown that halogen and carboxyl radicals may function either positively or negatively, depending principally upon the relative positions they may occupy in the benzene nucleus. Direct evidence that when intro groups occupy positively while the other functions negatively is shown by the replacement of only one of the nitro groups by negative hydroxyl or oxymethyl on interaction with Na. OH or Na. OCH, and the concomitant formation of Na. NO₂. For example, 1, 2 dintro- and 1, 2, 4-trinittobenzenes interact readily with solutions of sodium hydroxide or sodium methylate. The nitro group in

position 1 is replaced by OH or OCH₄, and Na. NO₂ is formed. These reactions find an interpretation in the electronic formula of benzene and may be summarized in the following equation —

Other facts which not only indicate the existence of negative uitro groups but also further substantiate the electronic formula of benzene are recorded in the recent work of P. van Romburgh and I. W. Wensink ** who found that the mto group in position 3 of cither 2, 3, 4 - 01 3, 4, 6-timutodimethylamline is readily replaced cither by an NH₄—on NH(CH₄)—

group on interaction with alcoholic solutions of ammonia or methylamme, respectively. Ordinarily, the hydrogen atoms of ammonia are positive and the three valences of the nitrogen atom are negative. Hence the abbieviated electronic formulae of

ammonia and methylamine are \vec{H} . \vec{NH}_2 and \vec{CH}_3 . \vec{NH}_3 . Accordingly, the replacement of only one of the three nitro groups

(that in position 3) by NH2 is perfectly analogous to the re-

placement of NO_2 by OH in the reactions given above. Furthermore, the nitro groups in positions 2 and 4 in the 2, 3, 4 compound are regarded as positive since they are not replaceable

by NH_2 They also occupy positions ortho to the replacable negative into group of position 3. Again, in the 2, 4, 6-compound the nitro groups in positions 4 and 6 are also regarded as

positive since they are not replaced by NH₂. They occupy positions ortho- and paia-, respectively, to the negative nitro group of position 3. All of these facts and conclusions are embodied in and interpreted by the following electronic formulæ and equations:—

In all benzene formulæ positions 2 and 6 are equivalent as are also positions 3 and 5. Accordingly, the 2, 3, 4- and 3, 4, 6-transformethylandines (and their products of interaction with ammonia) coincidentally show that when the substituents in positions 1, 3, and 5 are negative, the substituents in positions 2, 4, and 6 are positive. This also constitutes a further substantiation both of the electronic formula of benzene and the substitution rule.

Additional evidence leading to the above conclusion might be described in detail but buef mention may suffice. Recently Michele Giua 57 and A Contaidi 58 have described a number of polynitro derivatives of benzene which suffer replacement of a nitio group by OH, O(CH,), NH,, and NH(CH,) with the liberation of nitrites In some instances hydrolysis with water yields nitrous acid anhydride. In these various instances two nitro groups are adjacent, i.e., occupy positions ortho to each other, and only one of them suffers replacement. Giua maintains that this behaviour substantiates the rule on the reactivity of the nitro groups in the benzene ting proposed by Korner and Laubenheimer who attribute the facility of reaction of the one nitio group to the adjacence of the two nitro groups and the consequent "loosening" action of one group upon the other. This interpretation is not only vague but it fails to take into account the highly significant facts that the nitro group is replaced by a substituent of negative polarity, and that the group thus replaced is liberated as a nitrite in which the nitro radical is also negative. The electronic conception of positive and negative valences and the existence of positive and negative nitro groups affords a more satisfactory interpretation, since all of the facts noted are indicated and correlated by the electronic formula of benzene and the substitution rule. Conversely, the facts verify the electronic formula of benzene and the substitution rule

In concluding this chapter, it may be remarked that the electronic conception of positive and negative valences and the

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electronic formula of benzene afford explanations of many anomalous properties and teactions which are generally classed as stereochemical problems. The possibility of miterpreting some of these properties and teactions indicates that they are not necessarily anomalous or irregular. Hence it is hoped that a further extension of the principles and methods herewith presented may lead to the interpretation and solution of other problems.

CHAPTER IX

AN INTERPRETATION OF THE BROWN AND GIBSON RULE

No single subject in the history of modern chemistry has received more attention than that of substitution in the benzene nucleus. Concerning the introduction of two and three substituents, the foremost authority, A. F. Holleman, as states that he has found 1300 memoirs, and, if we include the formation of other polysubstituted derivatives, this number is increased to 2000. Many rules and hypotheses have been proposed to correlate and explain these substitution reactions. The foremost are those of Hubner, So Noelting, Armstrong, Brown and Gibson, Collie, Lapworth, Blanksma, Flurscheim, Obermiller, and Tschitschibabin. So

A comprehensive review of the rules and hypotheses of most of the foregoing investigations is given by Holleman in his work, Die direkte Einfuhung von Substituenten in den Renzelken (1910), and in his lecture before the Nociété Chimque de France (1911) entitled Sur let règlez de substitution dans le nayau benzénique. Despite the immense amount of work accomplished in this field, Holleman states that no rule or theory has as yet furnished a satisfactory explanation of substitution in the benzene nucleus. "The question of knowing the reason why such or such group directs a new substitution, is not yet solved."

Of all the rules that have been proposed for determining whether a given benzene monoderivative shall give a meta-diderivative or a mixture of ortho- and para-di-derivatives, that of Professor A. Crum Brown and Dr. John Gibson vis pne-eminent. Quotations from their original article will describe their rule

A. The Brown and Gibson Rule.

"When a monobenzene derivative C_bH_bX is so treated as to give a dibenzene derivative C_bH_bX , it is well known that, as a rule, this dibenzene derivative is either (a) exclusively, or nearly

so, a meta-compound, or (b) a mixture of ortho- and fara- with none or very little meta-, and that whether the case shall fall under (a) or (b) depends on the nature of the radical X, and not at all, or only very slightly, on the nature of Y.

"We have gone over all cases known to us, and have formulated a rule which holds in all these cases, and is always capable of rigorous application, so that, if in any case it should be found to fail, no excuse could be found foo it. We shall best explain our rule by showing how we apply it We shall write in column A a number of benzene mono-derivatives, in column B the radicles replacing one H of C₄H_m in column C the hydrogen compounds of these radicles, in column D the hydroxycompounds of the same radicles, and in column E the letters m or a-p according as the mono-derivatives give meta- or a mixture of or tho- and para. di-derivatives.

Λ	В,	С	D	Ŀ
C,H,CI C,H,Br C,H,CH, C,H,CH, C,H,NH, C,H,NO, C,H,CO, C,H,CO, C,H,CO, H C,H,CO, OH C,H,CO, OH C,H,CO, OH C,H,CO, OH	CI .Br .CH ₃ .NII ₂ .OH .NO ₂ .CCI ₃ .CO. H .CO. OH .SO ₂ .OH .CO. CH ₃ .CO. CH ₃	HCI* HBE* HCH.* HNH4.* HNH4.* HNO.* HCCI,* HCO.H H CO OH HSO.* HCC CH; HCC CH; HCC CH; HCC CH;	HOC! HOB; HOCH, HONH; HO. OH HO. NO;* HO. CCI, HO CO. OH * HO CO. OH * HO CO. CH;* HO. CH;. CO. OH	o-p o-p o-p o-p o-p m v-p m m m

"In column C, we mark with an asterisk those substances which are not capable, by direct oxidation, of being converted into the corresponding hydrogen compounds in the next column, and in column D we maik with an asterisk those substances which can be formed by direct oxidation from the corresponding hydrogen compounds so that in each horizontal line there is one asterisk and one only, either in column C or in column D. By direct oxidation we mean oxidation by one step. Thus, no doubt H. CH₃ can be converted into HO. CH₃, but not by one step of oxidation, whereas H. CO CH₄, can by one step of oxidation be converted into HO. CO CH₃. Now it will be seen that wherever the H compound is asterisked, we find σ in column E, and whenever the HO compound is asterisked, we find σ in find m in E, and whenever the HO compound is asterisked, we find σ in find m in m

column E. In other words, when X is naturally to be regarded as a derivative of HX, C, H, X gives ortho- and para-di-derivatives, and when X is naturally to be regarded as a derivative of HOX, C. H.X gives meta- di-derivatives Our test by which we determine whether X is to be regarded as derived from HX or HOX is, can HX be directly, by a single oxidizing step, be converted into HOX or not? If it cannot, then X is to be looked on as derived from HX, if it can, then X is to be looked on as derived from HOX"

Consider now the nitration of chlorobenzene and the chlorination of nitrobenzene in terms of the Brown and Gibson Rule, According to this rule C.H.Cl is looked upon as a derivative of H Cl which is not directly oxidizable to HO. Cl. Therefore C.H.Cl yields ortho- and para- chloronitrobenzene on nitration. On the other hand, CaH, NO2 is regarded as a derivative of H . NO, which is directly oxidizable to HO NO. Therefore, C.H. NO, yields meta-chloronitrobenzene on chlorination. Thus the experimental facts and rule are correlated. What is the underlying cause of this agreement between rule and fact? Brown and Gibson state that their rule is not a law, "because it has no visible relation to any mechanism by which the substitution is carried out in one way rather than in another way, but, if it is found to be rigorously applicable it must be related in some way to a law, and may be of use, both as grouping phenomena together and in guiding us to the cause of the formation of meta-compounds in certain cases and of ortho- and para- in others".

formula of benzene enable us to show that there is an underlying cause, a mechanism, so to speak, by which "substitution is carried out in one way rather than in another way," or, in other words, why some substitution reactions conform to the ortho-, para- type while others proceed according to the metatype. Both the reason and the mechanism become apparent when we determine the difference between X in HX, and X in HOX In our ordinary structural formulæ no difference is apparent, but from the electronic point of view, when H is positive and O is negatively bivalent, it necessarily follows that X in HX functions negatively while X in HOX functions positively as is evident in the electronic formulæ H--X and

The electronic conception of valence and the electronic

 $H^{+-}O^{-+}X$. Accordingly, a mono-substituted derivative may be either $C_e^{\dagger}H_s$, X_s , if it is regarded as a derivative of H. X_s or $C_e^{\dagger}H_s$, X_s if it is looked upon as a derivative of HO. X. Therefore (1) when X in $C_e^{\dagger}H_sX$ is negative and the entering substituent Y is positive, then ortho- and para-di-derivatives result. On the other hand (2) when X in $C_e^{\dagger}H_sX$ is positive and the entering substituent Y is positive, then meta-di-derivatives are found. These principles are embodied in the following typical equations:—

Thus, the electronic formula of benzene is correlated with the electronic interpretation of the Brown and Gibson rule, thereby showing a visible relation to a mechanism by which substitution is carried out in one way rather than in another way. Section A of the preceding Chapter VIII, presents concrete examples of the above scheme in the nitration of chlorobenzene,

C_kH₅Cl, and in the chlorination of nitrobenzene, C_kH₅NO₂. The electronic formula: of the products of these interactions, namely, ortho- and paia- and meta-chloronitrobenzene, are correlated not only with the methods of their preparation and their chemical properties, but also with the Brown and Gibson rule.

In many substitution reactions, ortho, para, and metadi-substituted derivatives of benzene are formed simultaneously. An interpretation of these simultaneous reactions necessitates a further development of the principle of electronic tautomerism

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which will be presented after a further analysis of the Brown and Gibson rule has been considered.

B. A Further Analysis of the Brown and Gibson Rule.

In presenting a further analysis of the Brown and Gibson unle it is necessary, first, to bring out more fully the hidden significance of the difference between compounds of the type HX which are not directly oxidizable to HOX and those which are directly oxidizable to HOX.

Some writers have taken exception to this point of view which Brown and Gibson assumed as their basis of classifying substituents For instance, Holleman 71 states that the rule "although remarkable and representing very well many of the observed facts, does not give satisfaction since it depends entuely upon the question of the direct oxidation of the hydrogen compounds of the substituents. Thus, at the time (1892) when Brown and Gibson published their rule, methane, H. CH, could not be exidized directly to methyl alcohol, HO, CH,, hence, according to the rule, the radical CH, should direct the entering substituent to the oitho and para positions. But since the time of the publication of the rule. Bone has proved that the pumary oxidation product of methane during combustion is methy alcohol. As a consequence, the radical CH, should then direct the entering substituent to the meta position. This discovery of Bone renders the rule inexact. While it is perfectly true that one cannot directly transform H Cl to HO, Cl, yet on the other hand no one is certain that this will always be impossible." Therefore, Holleman contends that with the progress of science the Brown and Gibson rule becomes less and less applicable.

Now this criticism of Holleman is only apparently justified, because Brown and Gibson unfortunately based their classification of substituents upon the debatable standpoint of the direct oxidizability in one step of HX to HOX. It has been shown that the chief point of distinction between the two classes of substitutions is not the direct oxidizability of HX to HOX but depends rather upon the electronic interpretation of the rule in conjunction with the electronic formula of benzene, and the fact that X in HX is negative while X in HOX is positive. These

points of view invalidate Holleman's criticism of the Brown and Gibson rule.

Further light may be thrown upon the nature of the substituents (X) commonly found in the benzene nucleus by considering the behaviour of certain HOX compounds as oxidizing agents, and the susceptibility to oxidation of certain HX compounds, thus placing the Brown and Gibson classification on a different basis Consider, first, a few of those substituents (OH, NH2, Cl, and CH2), which Brown and Gibson regarded as naturally derived from HX, not directly oxidizable to HOX, It is particularly noteworthy that the hydroxyl compounds of each of these substituents may function as an oxidizing agent. while their hydrogen compounds are not oxidizing agents in the common sense of the word. Furthermore, the hydroxyl compounds of these substituents are less stable than their corresponding hydrogen compounds. When HOX functions as an oxidizing agent (HOX -> HX + O), yielding HX and oxygen (either free or combined) there is a change in the polarity of X

from $\overset{\cdot}{X}$ to $\overset{\cdot}{X}$. This change may be represented as involving either an oxidation of negative hydroxyl ions by positive chlorine

ions (HO , Cl = HO + Cl \rightarrow H + Cl + O) or an intramolecular oxidation and reduction

 $(H\overset{+}{-}O\overset{+}{-}X \Rightarrow H\overset{+}{-}O\overset{+}{-}X \Rightarrow H\overset{+}{-}X + \overset{+}{0}) \quad \text{Now,}$ since compounds of the type HOX are less stable than those of the type HX, as shown by oxidation reactions, and since these

oxidations involve the transition from X to X, we may accord-

ingly assume that X is more stable than X. In other words, the tendency for X to function negatively is more pronounced than its tendency to function positively, but it may function

either way as its \dot{H} X and \dot{H} O \dot{X} compounds show. This may be po-fulated as a general property of those substituents which Brown and Gibson regarded as naturally derived from HX not directly oxidizable to HOX.

In order to complete the analysis of the Brown and Gibson rule it will be necessary to consider from the same points of view a few of those substituents, namely, CO₂H, COH, SO₃H

as O functions as the oxidizing agent through the acquisition of electrons and the consequent conversion of certain negative valences in the radical X to positive valences—

$$\stackrel{+}{0} \rightarrow \stackrel{-}{0} + 2 \oplus : \stackrel{-}{X} + 2 \oplus \rightarrow \stackrel{+}{X}.$$

The complete electronic equations for the oxidation of (1) formic acid, (2) formaldehyde, (3) sulphurous acid, and (4) nitrous acid are as follows:—

The above oxidations proceed readily, in fact most of the compounds are oxidized directly by atmospheric oxygen, $(O_z = O \xrightarrow{+} O \to \stackrel{+}{2} O)$. The tendency for the acid HX to change to HOX involves a change in the polarity of the complex radical X from \bar{X} to $\stackrel{+}{X}$, and accordingly $\stackrel{+}{X}$ may be regarded as more stable than \bar{X} . In other words, the tendency for X to function positively $(\stackrel{+}{HO} \cdot \bar{X})$ is more pronounced than its

tendency to function negatively (H, X), but it may function either way as the hydrogen and hydroxyl compounds show. The above may be postulated as a general property of those substituents which Brown and Gibson regarded as naturally derived from $\Pi(OX)$ which results from the direct oxidation of

Other points of significance may be noted in Equations 1-4 above. In formic acid, equation 1, three of the carbon valences are positive while the fourth is negative. In carbonic acid, four

HX

carbon valences are positive. Prior to the action of O, formic acid is in equilibrium with a tautomeric modification in which the carbon atom is bivalent and shows one free positive and one

free negative valence. These free valences unite with \overline{O} and, either simultaneously or subsequently, the negative carbon

valence is oxidized to a positive valence and oxygen (O) thereby

becomes negatively bivalent (O) The tautomeric modification of formic acid, showing a free posttive and a free negative valence, would through the polarization of these valences present an illustration of a bivalent carbon compound.

In this connection it should be noted that Stieglitz ⁷¹ has given a detailed account of the oxidation of carbon compounds, in particular formaldehyde, showing by means of the chemometer that there is involved an actual transfer of electrical charges. The classification of sniphurous and nitious acids with formaldehyde in the present discussion leads to the suggestion that certain sulphites and intrites might lend themselves to similar significant demonstrations.

In Equation 3 note that sulphurous acid (H. SO,H) is ordinarily written (HO. SO,H), in which the sulphur atom (comparable to the carbon atom above) shows a free positive and a free negative valence, and the polarization of these valences renders sulphur quadrivalent; but, on oxidation to sulphuric acid, the sulphur atom becomes sexavalent. In precisely the same manner intous acid (H. NO, is commonly written (HO. NO) in which nitrogen is tervalent, but oxidation to intic acid renders nitrogen quinquevalent. Now Equations 1-4 show that formaldehyle, formic acid, sulphurous acid, and nitrous acid

belong to the same class and same type of oxidizable compounds. Since the sulphu atom in sulphurous acid increases its valence by two on oxidation to sulphuric acid, and since the nitrogen atom in nitrous acid likewise increases its valence by two when oxidized to nitric acid, it follows, similarly, that the carbon atom in formaldehyde and formic acid must undergo likewise an increase of two in its valence when the compounds are oxidized to formic acid and carbonic acid, respectively. In other words, this classification of the substituents and the electronic system of oxidation lends further support to, and an electronic interpretation of, the existence of bivalent carbon compounds.

The extended interpretation and analysis of the Brown and Gibson rule brings out the following facts —

1. Certain sub-tituents (OH, Cl, B_1 , I, NH_2 , CH_2 , etc.) lead to the formation chiefly of ortho- and para-derivatives of benzene, because their tendency to function negatively is more pronounced than their tendency to function positively

This tendency may be formulated thus: $\ddot{X} > \ddot{X}$,

2. Certain other substituents (CO₂H, COH, SO₂H, NO₂, etc) lead to the formation chiefly of meta-derivatives of benzene, because their tendency to function positively is more pronounced than their tendency to function negatively. This tendency may be formulated thus: X > X.

3 The objections which have been disected against the Brown and Ghuson rule have been based upon the question of the direct oxidizability of HX to HOX. These objections are invalid, because the distinction between the two classes of substituents rests upon the facts summarized in paragraphs (1) and (2) above, and the fact that X in HX is negative while X in HOX is positive. These distinctions, in conjunction with the electronic formula of benzeus, show why certain mono-substituted derivatives of benzeus yield chiefly ortho- and para-di-substituted derivatives, while others yield chefly meta-fissibilitized derivatives.

CHAPTER X

ELECTRONIC TAUTOMERISM OF BENZENE DERIVATIVES

IN Section D of Chapter 11, and in Section C of Chapter 111, the conception of electronic tantomerism was developed. The purpose of the present chapter is to consider a number of experimental facts and to show that their interpretation not only fully wariants but also demands the assumption of the existence of electromers in dynamic equilibrium, i.e., electronic tantomerism. This conception was first proposed by the author in 1908. The following works "embody the definition:—

"The hypothesis of positive and regative valences suggests the possibility of there being at least two distinct types or electionies of a binary compound. Hence two mono-substituted

derivatives of benzene, $c_2H_3^+$ —X and $C_6H_8^-$ —X, should be possible while only one is known. Therefore, if both electroniers exist and only one substance is identified, it follows that the electroniers are either not distinguishable by (present) physical methods, or that one electronier is transformable into the other through the transposition of electrons:

$$C_aH_a \xrightarrow{+-} X \Leftrightarrow C_6H_bX \Leftrightarrow C_6H_6 \xrightarrow{--+} X$$
"

A study of various hydrolytic reactions compels one to assume the existence of such electromers. As previously noted, when a compound, XY, on hydrolysis yields HY and HOX,

then the compound is represented by the formula $X \cdot \overline{Y}$. The hydrolytic reaction conforms to the following scheme (a) —

If, on the other hand, the conditions of hydrolysis are such that compound XY yields HX and HOY, then XY must possess the

formula X . Y. The hydrolysis conforms to the scheme (b) --

$$(b)\stackrel{-}{X}\stackrel{+}{.}\stackrel{+}{Y} + \stackrel{+}{\text{II}}\stackrel{-}{.}\stackrel{-}{\text{OH}} \rightarrow \stackrel{+}{\text{II}}\stackrel{-}{.}\stackrel{\times}{X} + \stackrel{-}{\text{IIO}}\stackrel{+}{.}\stackrel{+}{Y},$$

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Now it has been found that phenylsulphonic acid, C_4H_8 . SO_2H , on hydrolysis, in *alkalme solution* yields sulphunous acid H. SO_3H , and phenol, C_6H_8 . OH. This reaction conforms to scheme (a) thus —

On the other hand, the hydrolysis of the same compound, phenylsulphonic acid, in acid solution yields benzeue, H C_iH₃, and sulphuric acid, HO. SO₃H. This reaction conforms to scheme (\$\text{\text{\$\e

These facts compel us to conclude that phenylsulphonic acid may enter into chemical reactions either as $C_c^{\dagger}H_s$, SO_4H or as

C_oH₃, SO_dH. From the tautomeric point of view, phenylsulphonic acid may be regarded as an equilibrium mixture of two electronic isomers, or electromers

$$(C_0\overset{+}{H}_5.S\overset{-}{O_3}H \Rightarrow C_0\overset{-}{H}_5.S\overset{+}{O_3}H)$$

and any other mono- or poly-substituted derivative of benzene may be regarded similarly, as will be shown later

Consider any mono-substituted derivative of the general formula CaHaX which will now be regarded as

$$(C_6\overset{+}{H}_5\cdot \widetilde{X} \iff C_6\overset{-}{H}_5\overset{+}{X}).$$

Indicating only the polarities of the hydrogen atoms of the nucleus, the equilibrium is represented thus --

Note particularly that the change in the polarity of the substituent X involves simultaneously a change in the polarities of the hydrogen atoms, but that in each electromer those atoms (or substituents) ortho and para to X are opposite in sign to X, while those atoms (or substituents) meta to X are of the same sign as X. This, of course, is the essential feature of all of the electronic formulae of benzene, namely, that hydrogen atoms or substituents in positions 1, 3, and 5 of any given electromer are opposite in sign to the hydrogen atoms or substituents in positions 2, 4, and 6 of the same electromer, otherwise the symmetry, and consequently the stability, of the benzene uncleus could not be maintained. A detailed mechanism of the transposition of electronic hardward electronic relative valences, etc., involved in the electronic tautomerism of benzene derivatives will be given later. For the present, the simpler formulae involving only the hydrogen atoms and substituents are sufficient. In a similar manner the electronic tautomerism of a dissubstituted derivative of benzene, for example C₂H₂XY₂ in which X and Y are in para positions to each other, may be propresented this. —

When two substituents occupy positions either ortho or para to each other they are of opposite polanty. Hence in the electronic formula for a p-dihalogen between in which the halogen atoms X and Y are different chemically, it follows that if X is negative, Y is positive, o, if X is positive of year and intration of a p-disabstituted derivative of benzene only two mono-initio substitution products are possible, and, from the standpoint of the principle o' electronic tautomerism as applied to a p-dihalogen benzene, the nutation reaction should conform to the following general scheme:—

If the principles underlying this general scheme are correct, then certain definite consequences follow which should lend themselves to verification by experiment. What are some of these consequences? If the electromers of C,H,XY (designated as A and B) are in

tautomeric equilibrium and if each electromer interacts with nitric, agid, the respective isomers A' and B', should be formed. Note particularly that in the isomer A', X is negative and Y is negative. In view of the facts previously presented, notably the action of sodium hydroxide or sodium methylate upon \(\text{\sigma}, \text{\gamma}, \text{\gamma} \), and \(\text{\gamma}, \text{\text{\gamma}} \) and the chloronitrobenizenes, it follows that isomer A' on interaction with sodium methylate (Na. OCH.) should exchange X for OCH, and

sodium methylate (Na. OCH₂) should exchange X for OCH₃ and liberate Na. X according to equation (1) -

$$(t) \left(\begin{matrix} \widetilde{X} \\ A' \end{matrix} \right)^{NO_2} + \begin{matrix} \downarrow \\ N_A, \widetilde{OCH}_2 \end{matrix} \rightarrow \left(\begin{matrix} \widetilde{OCH}_2 \\ A'' \end{matrix} \right)^{NO_2} + \begin{matrix} \downarrow \\ N_A, \widetilde{X}. \end{matrix}$$

On the other hand, isomer B' should exchange \hat{Y} for OCH₈ and liberate Na. \hat{Y} according to equation (2):—

(a)
$$\begin{pmatrix} \dot{x} \\ B' \\ NO_2 \end{pmatrix}$$
 + $\begin{pmatrix} \dot{x} \\ Na \\ OCH_1 \end{pmatrix}$ $\Rightarrow \begin{pmatrix} \dot{x} \\ B'' \\ NO_2 \end{pmatrix}$ + $\begin{pmatrix} \dot{x} \\ Na \\ V \end{pmatrix}$

Furthermore, if electromers A and B are in tautomeric equilibrium and nutration yields a muxture of isomers A' and B' then the action of sodium methylate upon this mixture should yield not only the oxymethyl derivatives, A'' and B'' (equations (1) and

(2), respectively), but also a mixture of the sodium haldes, Na. X

and $\stackrel{\frown}{Na}$. An experimental verification of these predictions would lead to a very definite conclusion, namely, that the paramixed dishalogen compound, C_6H_1XY , affords an example of electronic landomerism. Otherwise it would be difficult, if not impossible, to account for the simultaneous formation of the mixed

halides, Na. X and Na. Y, when sodium methylate interacts with the products of nitration of the mixed dihalogen benzene.

Experimental facts will now be considered which are completely in agreement with the principles and general reactions just postulated. Holleman and Heineken have made a thorough study of the nitration (at 0°) of various mixed diladogen henzenes. For the present, attention will be confined to the intration of p-chlorobromobenzene, which gave a nitxture of the isomers 1, 2, 4- and 1, 3, 4-chlorontrobromobenzenes. In developing a quantitative chemical method for determining the yields of these somers in the nitration mixture, Holleman found that the 1, 2, 4-isomer heated with 0.35 N sodium methylate solution in a sealed tube for five hours at 80° gave practically complete displacement of chlorine as solium chlorate while the bromum remained intact. The reaction proceeded quantitatively (68.50 piez cent) according to equation (3):—

(5)
$$A \mid NO_2 + Na, OCII_d \rightarrow A'' \mid NO_2 + Na, CI$$

$$\bigcup_{Dr} (t_1, t_2, t_3)$$

Note that this reaction is perfectly analogous to equation (1) given above, in which \widetilde{X} becomes \widetilde{Cl} , and \widetilde{Y} becomes \widetilde{Bi} . The liberated sodium halide is \widetilde{Na} . \widetilde{Cl} . In other words, the 1, 2, 4-isomer (\widetilde{A}) has the electronic formula

Otherwise the liberation of chlorine as Na Cl is mexplicable,

On the other hand, the 1, 3, 4-somen on interaction with sodium methylate (under dentucal conditions) suffered practically complete displacement of bromine as sodium brounde while the chlorine atom remained intact. The reaction proceeded quantitatively (98 vo per cent), according to equation (4).

(4)
$$\begin{array}{c} CI \\ B' \\ NO_3 \\ Bc \\ (t, 3, 4^2) \end{array}$$
 + Na OCH₃ \Rightarrow $\begin{array}{c} CI \\ B' \\ OCH_3 \end{array}$ + Na Be

Note that this reaction is perfectly analogous to equation (2) above, in which $\stackrel{.}{X}$ becomes $\stackrel{.}{Cl}$, and $\stackrel{.}{Y}$ is $\stackrel{.}{Br}$. The liberated sodium halide is $\stackrel{.}{Na}$ $\stackrel{.}{Br}$. In other words, the I, 3, 4-isomer (B) has the electronic formula



In no other way can the elimination of bromine as Na . Br be explained.

The quantitative interaction of 1, 2, 4- and 1, 3, 4-chloro-introbromobenzene with sodium methylate enabled Holleman to determine the relative quantities of these isomers obtained on ritiating f-chlorobromobenzene. The sodium chloride and sodium bromde liberated were precipitated as silver chlorobrome and silver bromide. An analysis of the mixed precipitates gave the percentage yields of the isomers calculated upon the ratios of AgCl: 1, 2, 4-isomer, and AgBr. 1, 3, 4-isomer. These calculations were based upon equations (3) and (4) which are perfectly analogous to the preceding electronic equations (1) and (2), respectively. The yields of the isomers were 45 20 per cent and 5,480 per cent, respectively. The complete correlation of all of these reactions with the electronic formulae and the principle of electronic tautomersm is embodied in the following scheme —

Now a survey of the above scheme shows that those molecules of I. 4-chlorobromobenzene which interact with nitric acid to give the 1, 2, 4-isomer, which in turn exchanges its chloring atom for OCH, and yields sodium chloride, cannot be identical with the molecules of 1, 4-chlorobiomobenzene which interact with nitric acid to give the 1, 3, 4-isomei, which in turn exchanges its bromine atom for OCH2 and yields sodium bromide, In other words, the 1, 2, 4-isomer (A'), containing Cl and Bi must be derived from molecules of 1, 4-chlorobromobenzene which also contain Cl and Br. i.e. electromer A. and the 1. 3. 4-isomer (B') containing Cl and Br. must be derived from molecules of 1, 4-chlorobromobenzene which also contain Cl and Br. i.e., electrome: B. Thus the existence of two types of molecules of 1, 4-chlorobromobenzene, namely, electromers A and B, is conclusively indicated. Their simultaneous interaction with artric acid clearly indicates tautomeric equilibrium. The relative yields of the isomers A' and B' are functions of the speeds of interaction of nitric acid with the respective electromass A and B, and, as will be shown in a subsequent chapter, the speeds of interaction are intimately related to the concentrations of the respective electromers in the reaction mixture.

It will not be necessary to extend this chapter by giving similarly detailed descriptions of the nitrations of other mixed dihalogen benzenes, such as p-fluorochloro. p-fluoroiodono, p-chloroiodon, p-chloroiodon, p-chloroiodon, p-chloroiodon, p-chloroiodon, and p-bromoiodo benzene Each example conforms perfectly to the schemes and principles just presented In conclusion, it is maintained that each of these cases further substantiates the principle of electronic tautomerism or the existence of electronics in dynamic equilibrium.

Any objection to the use of the conception of electronic tautomerism on the grounds that electronic isomers (electromers) have not as yet been isolated is unwarranted because the conception of tautomerism, both in its historical and practical aspects. neither involves nor demands the immediate possibility of the isolation of the tautomeis. In 1885, Laar, 76 in a paper Ueber die Moelichkeit mehrerer Strukturformeln für dieselbe chemische Verbindung, defined tautomerism as the phenomenon of one substance doing duty for two structural isomers, but the isolation of dynamic isomers or desmotrores was not effected until 1803, the earliest examples being acetyldibenzovlmethane, tribenzovlmethane, and mesityloxide-oxalic ester described by Claisen.76 Strictly speaking, the term tautomeric is now applied to substances in which only the eauthbrium mixture of the structural isomers is known: the term desmotropic is employed in cases where the isomers have been shown to be capable of independent existence.77 The existence of electromers in dynamic equilibrium has been conclusively indicated. Their isolation as electronic desmotrones has not as yet been effected. Further discussion of this phase of the subject is considered in Chapters XXIII, and XXIV.

The phenomenon of electronic tautomensm is met repeatedly in the interpretations of substitution reactions in the benzene nucleus. A number of these reactions is described in the following chapter

CHAPTER XL

THE SIMULTANEOUS FORMATION OF ORTHO-, PARA-, AND META-SUBSTITUTED DERIVATIVES OF BENZENE

A. Formation of Isomeric Di-substituted Derivatives.

In the electronic interpretation of the Brown and Gibson rule (Chapter IX.), the formations of ortho- and para-compounds on the one hand, and meta-compounds on the other, were regarded as distinct and separate processes because, as a general rule. a given mono-substituted compound yields chiefly a meta-diderivative or a mixture of ortho- and para-di-derivatives. In many substitution reactions isometic ortho-, para-, and meta-disubstituted derivatives are formed simultaneously. The Brown and Gibson jule does not cover these cases. Consequently it becomes necessary to explain (A) the simultaneous formation of isomeric ortho-, para-, and meta-di-substituted derivatives, and also (B) the simultaneous formation of isomeric poly-substituted derivatives of benzene. These explanations involve the electronic formula of benzene, the rule of substitution, and the conception of electronic isomers or e'ectromers in dynamic equilibrium, 1 e., electronic tautomerism

It is necessary, first, to recall a well-established fact, namely, that in a mixture of two isomes (A = B) capable of changing one into the other with such readmess that the balance between them is rapidly restored, the addition of a given substance C will result in reaction either with A, or with B, or with both A and B. If interaction occurs between A and C, B will be transformed into A on account of the disturbance of the equilibrium. The mixture then reacts as though it consisted wholly of A. If C reacts only with B, the mixture behaves, for the same reason, as though B were the only substance present. In some instances two simultaneous independent reactions may occur, namely, the mixture behaves for the same reason of A and C simultaneously with the interaction of B

and C. In a precisely similar manner a given reagent, HO. Y, in the presence of a tautomeric mixture of electromers

may react with the one electromer or the other, or with both electromers. A complete scheme of interaction follows —

It should be observed that if \vec{HO} . $\vec{\hat{Y}}$ reacts only with $C_n^{\dagger}H_n$. \vec{X} (electromer A) then only ortho- and para-di-substituted

 $C_6\Pi_6$. A (electroner A) then only of the and para-di-substituted derivatives (A' and A'') are formed. Such is the case when chlorobenzene is intrated: the products of the interaction are

ortho- and paia-chloronitrobenzene. If \overrightarrow{HO} , \overrightarrow{Y} reacts only with C_6H_a , \overrightarrow{X} (electromer B), then only a meta-di-substituted derivative (B) results. Such is the case when nitrobenzene is chlorianted the product of the reaction is meta-chloronitro-

benzene. If \overrightarrow{HO} , \overrightarrow{V} interacts simultaneously with both electromers (A and B) then ortho-, meta-, and para-di-substituted derivatives (A', B', and A'', respectively) will be formed. The possibility and the extent of these reactions (as in all reactions with tautometic substances) are dependent upon several conditions such as the solvent medium, catalytic agent, temperature, concentration of the electromers, the speed of transition of one electromer into the other, and the respective speeds of interaction of the substituting agent with the respective electromers. These conditions are the problems which confront us in all research

involving tautomers, and likewise will be met in the case of electronic tautomers

Having given a general scheme for the simultaneous formation of ortho, meta, and para-th-substituted derivatives, one of the many examples which are in accord with and, therefore, confirm the scheme, should now be considered. For instance, in the intration of toluene, the substituent (X) already present in the nucleus is the methyl radical which may function either positively or negatively, but in Section B of Chapter IX, it has been shown that its tendency to function negatively is more pronounced than its tendency to function positively, i.e.,

CH₃ > CH₃ Hence, in the tautomeric equilibrium-

the former electromer would predominate and interaction with

 \overrightarrow{HO} \overrightarrow{NO}_2 would lead to a greater yield of a- and f-introtoluenes than of m-nitrotoluene. The predominance or preponderance of

the electromer C_iH_2 , CH_2 over the electronic tautomer C_iH_2 , CH_3 , may be correlated with the speeds of reaction with nitrie acid, i.e., if the former electromer reacts more rapidly than does the latter electromer, then the yield of ortho- and para-nitrotolucies would exceed that of meta-nitrotolucies.

The predominance of one electromer over the other may be regarded as equivalent to a greater concentration of the one electromer. Since concentration is a principal factor of the velocity of reaction, it is evident that a direct relationship exists between the predominance or preponderance of a given electromer and the velocity of the substitution reaction in which it is engaged

The foregoing assumptions are confirmed by experimental facts, since toluene when nituated yields 56'00 per cent of the α_1 , 40 90 per cent of the p_1 , and 3'10 per cent of the m-nitrotoluene. The simultaneous formation of the three isomers is indicated in the following scheme —

Here electromer A predominates, i.e., its speed of interaction with nitric acid is greater than the speed of interaction of electromer B with nitric acid.

Consider now, on the other hand, the nitration of benzoic cidd. The substituent (X) already present in the nucleus is CO_2H , which may function either positively or negatively, but $CO_2^{\dagger}H > CO_2H$, i.e., the tendency for the carboxyl radical to function positively is more pronounced than is its tendency to function negatively. Accordingly, in the equilibrium—

$$C_aH_a$$
. $CO_aH \Leftrightarrow C_aH_a$. CO_aH ,

the latter electromer (in which the carboxyl radical is positive) would predominate, and if its speed of interaction with $\overline{\text{HO}}$. $\overset{\bullet}{\text{NO}_2}$ is greater than the speed of interaction between $C_e^{\dagger}\overline{\text{H}}_3$. $C_2^{\dagger}\overline{\text{H}}_3$ and $\overset{\bullet}{\text{HO}}$. $\overset{\bullet}{\text{NO}_2}$, then the yield of *m*-nitrobenzoic acid should exceed that of the combined yields of the ortho- and paracompounds. These assumptions are confirmed by the facts, since benzoic acid on nitration yields 80 30 per cent of the metacompound, 18 50 per cent. of the ortho, and 1 20 per cent. of the p-nitrobenzoic acid. The electronic equations conform to the general scheme given above (p. 94).

Many other examples of the simultaneous formation of orthometa-, and para-di-substituted derivatives of benzene might be presented in detail. They are in complete agreement with, and accordingly confirm, the principle of the electronic tantomerism* of the electronics of the benzene demantives, the general rule for substitution, and the scheme for the simultaneous formation of orthos, meta, and para-fa-substituted definitatives of benzene.

B. Formation of Isomeric Poly-substituted Derivatives.

The principles underlying the simultaneous formation of poly-substituted derivatives of benzene are identical with those just given in the preceding section (A) and may be illustrated by considering the introduction of a third substituent (Z) into (1) an ortho-, (2) a para-, and (3) a meta-di-substituted derivative, CHLXY.

Scheme (1)—Introduction of a Third Substituent into an Orthodi-substituted Derivative.

Here the electronic tautomerism involves two electronics of an ortho-di-derivative. These electronics will be designated by the letters A and B while their substitution products will be indicated by A', A'', and B', B'', respectively. The general scheme (1) for the introduction of the third substituent Z is as follows:—

Theoretically and electronically four tri-substituted isomers are obtainable, namely, A' and A'' from electronic A; and B'' and B'' from electronic B. A remarkable experimental confirmation of this scheme is found in the work of Wibaut^{1,6} who showed that the nitration at o' of o-chlorotoluene yielded all four possible mono-nitro-compounds in considerable quantity The electronic interpretation is embodied in the following equations, which are perfectly comparable to the typical equations of scheme (1) above The percentage yield of each isomer is indicated beneath its appropriate electronic formula —

It has been shown that CI>CI, and CH4>CH2, i.e., the chlorine and methyl radicals may tend in general to function negatively rather than positively, but in o-chlorotoluene (either electromer A or B) the substituents must be opposite in polarity. This leads to the consideration of a significant question, namely, will the polarities of the substituents present in the nucleus predetermine in any way the relative quantities of the isomeric derivatives which are formed? In the simultaneous formation of the isomeric di-substituted derivatives of benzene the polarities of the substituents and the tendencies of certain substituents to function in one way rather than in another way were shown to be related to the relative quantities of the isomers (ortho and para on the one hand, and meta on the other) which were formed. Now in the case of o-chlorotoluene the two substituents present naturally tend to function negatively, but, since one must function negatively while the other functions positively, another perplexing question arises, namely, is the tendency for chlorine to function negatively more pronounced than the tendency for methyl to function negatively? An answer to this important question can be approached only in an indirect way by comparing the general stability of methyl alcohol (HO. CH3) with that of hypochlorous acid (HO . Cl). The latter is undoubtedly the less stable, since it is readily decomposed by light or heat, whereas methyl alcohol is a comparatively stable compound. Now, the instability and decomposition of hypochlorous acid involves the change, as

previously shown, from Cl to Cl; and since HO. Cl is less stable

than HO CH., naturally it may be assumed that the tendency for chloring to function negatively is more pronounced than the

tendency for methyl to function negatively, i.e., Cl>CH, in equivalent terms, the tendency for methyl to function positively is more pronounced than the tendency for chlorine to

function positively, i.e., CH2 > Cl. (The relative tendency for a radical to function in one way rather than in another way may be defined as its polar stability.)

These relationships lead to the conclusion that of the two electromers (A and B) of o-chlorotoluene, the one (A), in which (CI) is negative and (CH2) is positive, would predominate, which in turn would indicate that the nitration of e chlorotoluene might yield a greater quantity of the nitiochlorotolucies A' and A", than of the nitrochlorotoluenes B' and B". These assumptions are confirmed by the facts, since Wibant (loc cit.) has determined the yields of the four mono-nitro-o-chlorotoluenes to which are assigned, herewith, the abbreviated electronic formulæ in conformity with the preeeding scheme of substitution :-

The combined yields of A' and A" total 62.2 per cent, as against 37'7 per cent., the combined yields of B' and B".

Attention should be called to the fact that the percentage yields of the isomeric products of nitration of ortho-chlorotoluene, in the preceding scheme (1), and also the percentage yields 100

of the isomeric products of nitration of para- and metachlorotoluene in the following schemes (2) and (3) respectively, were determined by physical methods—processes of finctional crystallization and compasison of curves of fusion of mixtures of unknown composition with those of known composition.

Scheme (2)—Introduction of a Third Substituent into a Paradi-substituted Derivative.

Consider now the introduction of a third substituent into a para-di-substituted derivative of benzene according to the following scheme (2),—

Theoretically and electronically, two and only two tri-substituted delivatives are possible, namely, A' and B' from electromers A and B respectively. One of several series of experiments which confirm the above scheme is the nitration at o° of photorotolene by Holleman, be who demonstrated that only two mono-nitio-p-chlorotoluenes were produced. The correlation of these facts with the electronic formulæ is embodied in the following scheme:

$$\left\{ \begin{array}{c} \overset{\overset{\circ}{\text{Cl}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\overset{\circ}{\text{H}}}}{\underset{\overset{\circ}{\text{H}}}}}{\underset{\overset{\overset{\overset}{\text{H}}}}}{\underset{\overset{\overset{\overset{\overset{\overset}{\text{H}}}}}}{\underset{\overset{\overset{\overset{\overset{\overset{\overset{\overset{\overset}{\text{H$$

Since $\vec{Cl} > \vec{CH}_3$, or $\vec{CH}_3 > \vec{Cl}$, one would be disposed to assume that electromer A would predomin the and, accordingly, the intration of para-chlorotoluene would be expected to yield a greater quantity of A', C_8H_2 , \vec{CH}_3 , \vec{Cl} , $N\vec{O}_2$ (t, 4, 3), than of B', C_8H_3 , \vec{Cl} , \vec{NO}_2 (t, 4, 2)

This assumption is not realized single the yields are 42'00 per cent of A' and 58'00 per cent. of

B. This leads to the assumption that the relative polar stabilities of substituents in para position to each other may differ from the relative polar stabilities of the same substituents when in the ortho position to each other. For example, in the preceding scheme (1) for the untration of ortho-chlorotoluene, electromer A predominates, but in scheme (2) for the initiation of para-chlorotoluene, electromer B predominates. Hence, in scheme (2) the speed of interaction of electromer B with intricacid is greater than the speed of interaction of electromer a with the substituting reagents determines the yields of the substituted products.

Scheme (3)-Introduction of a Third Substituent into a Metadissubstituted Derivative

The introduction of a third substituent into a meta-di-substituted derivative of benzene corresponds to the following general scheme:—

$$\begin{bmatrix} \bar{X} & \bar{X} & \bar{X} & \bar{X} \\ \bar{\Pi} & \bar{X} & \bar{Y} & \bar{\Pi} & \bar{\Pi} \\ \bar{\Pi} & \bar{A} & \bar{Y} & \bar{H} & \bar{\Pi} \\ \bar{\Pi} & \bar{A} & \bar{Y} & \bar{H} & \bar{\Pi} \\ \bar{\Pi} & \bar{A} & \bar{Y} & \bar{H} & \bar{\Pi} \\ \bar{\Pi} & \bar{A} & \bar{Y} & \bar{H} & \bar{\Pi} \\ \bar{X} & \bar{Y} & \bar{H} & \bar{H} \\ \bar{X} & \bar{X} & \bar{X} & \bar{X} \\ \bar{X} & \bar{X} & \bar{X} & \bar{X} & \bar{X}$$

Since the substituents X and Y occupy positions meta to each other they must be of the same polarity, i.e., both are negative in A while both are positive in B. Some interesting theoretical and experimental results pissent themselves, depending upon the tendencies of X and Y to function positively or negatively. If both X and Y belong to that class of substituents which lead to the formation chiefly of ortho- and para-derivatives

(which is the case when $\hat{X} > \hat{X}$ and $\hat{Y} > \hat{Y}$), then electromes A

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would predominate and the chief substitution products would be 'A', A', and A'''. Furthermore, since X and Y are meta to each other and each is naturally negative, electromer A might predominate to the exclusion of electromer B; and the introduction

of a third substituent \dot{Z} would lead to the formation of only hree isomeric tri-substitution products, namely, A', A'', and A'''Just such a case is found in the nituation at O' of m-chloro-toluene by Wibaut (loc. cit.) The percentage yields of the isomers are indicated in the scheme:—

Now it has been shown in some instances that the tendency for chloring to function negatively is more pronounced than the tendency for methyl to function negatively, i.e., C1>CH. leads to the assumption that the entering positive nitio radical would be more subject to the directing influence of Cl than of CH. Now Cl directs NO. to the para position rather than to the ortho position, since chlorobenzene on nitration yields 60.00 per cent, of p-nitrochlorobenzene, and 30 10 per cent, of p-nitrochlorobenzene. On the other hand, CH, directs NO, to the outho position rather than to the para positions since toluene on nitration yields 56:00 per cent, of the o- and 40:90 per cent, of the p-nitrotoluene. Accordingly, when p-chlorotoluene is nitiated, one should expect the greatest yield to be that of isomer (A') in which NO, occupies a position para to Cl and ortho to CH3. A smaller yield would be predicted for that isomes (A") in which NO, is ortho to Cl and para to CH. The smallest yield would be predicted for the isomer (A"') in which NOz is ortho to Cl and ortho to CH2. These assumptions are fully confirmed by the facts since the respective yields of the three somers, Λ' , Λ' , and Λ'' , are respectively, 58 or per cent, 2000 per cent, and 8 80 per cent. This constitutes a further proof that the polarities of the substituents in the nucleus and their tendencies to function in one way rather than in another way predetermine not only the type of "ubstritties" but also the

relative quantities of the isomers produced.

Returning again to the general scheme (3) above, it should be noted that if X and Y belong to that class of substituents which lead to the formation chiefly of meta derivatives, which is

the case when $\overset{\circ}{X}>\overset{\circ}{X}$ and $\overset{\circ}{Y}>\overset{\circ}{Y}$, then electronus B would predominate. Acro dingly, the intro-hector of a third substituent Z would lead to the for mation theigh of the tris-substituent product in which all three substituents (X, Y, and Z) are positive Many examples are found in the literature which show that when the substituents in the nucleus are positive, the entering substituent assumes a position meta to those prevent, and the predominating isomer corresponds to B, the derivative of electromer B Exact quantitative data upon the amounts of the other possible somers have not been found in the literature.

Having considered all of the possibilities which might be encountered in the introduction of a third substituent Z into an ortho, a pain, and a meta-di-substituted derivative, C,H,XY, of benzene, and having shown that experimental facts and data are in agreement with the theoretical deductions, with the electrone formulae, and the principle of electronic tautomenism, and with the general electronic rule for substitution, it should be added that the formation of other poly-substituted derivatives of benzene will conform in general to the principles and rule previously developed and illustrated. This claim is warranted by a careful study of the numerous tables of the variously substituted derivatives of benzene which have been compiled with great care and presented by Holleman (loc cit) in his extended work on the influence of the substituents in the benzene nucleus, and the principles of substitution

CHAPTER XII.

THE ACTION OF SODIUM METHYLATE ON THE PRODUCTS OF NITRATION OF THE ORTHO-, PARA-, AND META-CHLORO-TOLUENES.

In the preceding chapter it was shown that unless orthopara, and mate-shlorotoluem amiliest the phenomenon of electionic tautomerism it would be impossible to explain the simultaneous formation of the variously substituted isomerinto-derivatives when the chlorotoluenes are nitrated. The principle of electionic tautomerism involved in the explanation of these substitution reactions will receive further experimental confiunation if it is possible to show by hydrolytic reactions that some of the nitro-derivatives are the products of the interaction of one of the electionness of a chlorotoluene while the other isomeric nitro-derivatives simultaneously formed are the products of matation of the other electromer of the chlorotoluene.

To this end, attention must be directed throughout this chapter to schemes (1), (2), and (3) of the preceding chapter for the intrations respectively of (1) ortho, (2) paia, and (3) metachloronitobenzene. These schemes constituted the electronic explanation of the nitration reactions conducted by Wibaut, Holleman and Wibaut, respectively, who determined the perentage yields of the several isomers of each nitration by physical methods—processes of fractional crystallization and comparisons of curves of fusion of mixtures of unknown composition with those of known composition.

In each of these schemes it will be observed that the electionic formulæ of the isomeric nitration products (A', A", etc) of the electromer A embody negative chlorine while those (B', B", etc.) of electromer B embody positive chlorine. Now, if it be possible to show by quantitative chemical methods that the yields of the isomers (chloronitrotoluenes) which contain negative chlorine are commensurate with the yields of the same isomeric ascertamed by Holleman and Wibaut by physical methods, then the conception of the electronic tautomers in of the chlorotolucnes as indicated in schemes 1, 2, and 3 is, to this extent, further substantiated.

The chemical method for the determination of the yields of the isomers containing negative chlorine depends, first, upon the completeness of their interaction with sodium methylate and, second, the non-interaction of the isomers containing positive chlorine. In order to test the applicability of these two points, the action of sodium methylate in methyl alcohol solution upon ϕ , ϕ , and m-chlorontrobenzenes was investigated. These isomers were chosen because it is commonly recorded that the ϕ - and ρ -isomers exchange their halogen atom for OH, O(CH₃), or NH₂, while the m-isomer is non-reactive.

An interpretation of these reactions has been given in Chapter VIII., Section A, from the standpoint of the electronic formula of benzene and the substitution rule. The abbreviated electronic formulas of the chloronitrobenzenes are as follows—

Only those isomers containing negative halogen (o- and f-) should interact with sodium methylate, thus

$$NO_2$$
 C_0H_4 , C_1 + NA , OCH_1 $\rightarrow NO_2$, C_0H_4 , OCH_1 + NA , CI_4 and the *m*-isomet, containing positive chlorine, should be non-reactive under the same conditions.

The extent of the replacement of negative chlorine can be found by determining the quantity of sodium chloride liberated. After repeated experiments, designed to secure the conditions noted, the following method was found to yield theoretical results. About 0.2 gram of the chloroutrobenzene was dissolved in 15 cc. of a nounal solution of sodium methylate in absolute methyl alcohol. The solution was heated at 100° in a sealed glass tube for 5 hours. The contents of the tubes were then diluted to a volume of 200 c.c., acidified with nitric acd, bolled to expel the methyl alcohol, cooled, and filtered. The quantity of sodium chloride in the respective filtrates was determined by

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titration with standard solution of silver nitrate.	The following
results were obtained :	

Substance.	(Grums)	No cc o 9957N AgNO : Solution	Per Cent Negruve Cl 1 ound	Per Cent Negative Cl Theoretical
0.C1.C,H4.NO3	0 2005	13.81	22.56	22 5t
/ C1. C.H. NO.	0'2007	12.78	22:48	22 5T
m-Cl. C,H, NO,	0.1903	0.00	0000	0 00

These results show conclusively that isomers containing negative chlorine may be estimated quantitatively, provided conditions are secured for their complete interaction with sodium methylate, while under the same conditions the isomer containing positive chlorine is non-reactive

The chief problem of this investigation was to secure conditions under which the negative halogen atoms of the nitration products of o-, b-, and m-chlorotoluenes would completely interact with sodium methylate and thus afford a chemical method for determining the percentage yields of the isomers of chloronitiotoluene which contain negative chlorine. A comparison of the chemically determined and the physically determined yields could then be made With this end in view, o-, p-, and m-chlorotoluenes were each separately nitrated (at o°) according to the specific directions given by Wibaut, Holleman and Wibaut, respectively. The methods were originally designed to yield only mono-nitio substitution products. The respective products of nitration of each of the chlorotoluenes, which consisted of mixtures of the different isomers (nitrochlorotoluenes), as indicated in the preceding schemes I, 2, and 3, were treated with sodium methylate solutions of varying concentrations-0.5 N. N. 2 N. and 3 N. Samples, 0.2 to 0.4 gram, of the nitration products were heated, each with 15 c.c. of the sodium methylate solution in sealed tubes, for 10 hours at 100°. The contents of the tubes were then analysed, as previously described, for the

negative chlorine, liberated as Na. Cl, precipitated and weighed as silver chloride. The percentage yield of the combined isomers containing negative chlorine was calculated from the ratio

Ag Cl : Cl . NO. CH, C.H. The results of these experiments are recorded in three sections (I., II., and III.) to correspond with schemes 1, 2, and 3 for the nitration of e-, f-, and mchlorotoluenes, respectively. All reactions were conducted in duplicate

TABLE I. I ACTION OF SODIUM METHYLATE OR THE ISOMERS DERIVED TROM O CHI ORO LOLULN . (See Scheme 1)

		,		
Mixture of Isomers (gram)	Normality NaUCH ₂	AgCl (gram)	Per Cent Suld of Itomers V + A" (1, 2, 6 and 1, 2, 4 Cl, Cl13, No., CaHa)	Average
0°2625 0 2434 0 2835 0 2938 0°3283 0 2994 0°2772 0°2940	0°5 N 0°5 N N N 2 N 2 N 3 N 3 N	0 1163 0 1084 0 1362 0 1362 0 1446 0 1362 0 0941	51'ut 51'29 50'28 50'29 40'04 90 12 38'16 38'16	53°15 55'73 19'88 38'43

These results indicate that the extent of the reaction with sodium methylate is greatest in the normal solution. A 58 73 per cent, yield of the combined isomers containing negative chlorine was obtained. The yields of these isomers determined by the physical methods of Wibaut total 62:20 per cent. The yields obtained by the two methods, while not identical, approximate closely. Variations will be considered later.

TABLE IL II ACTION OF SODIUM METHYLAGE ON THE ISOMPRS DERIVED FROM A-CHLOROLOLULME (See Scheme 2)

Mixture of Isomirs (gram)	Normality NaOCII _d	AgC1 (gram).	Per Cent Yield of Isomer A'	Average
0.4707 0.4868 0.3668 0.2668 0.2658 0.3658 0.3068 0.3297 0.3037	0 5 N 0 5 N N N 2 N 2 N 2 N 3 N 3 N	0 0799 0 0842 0 0946 0 0773 0 0750 0 0780 0 0814 0 0769	35 33 35 13 36 89 35 17 31 11 30 42 29 54 30 730	35 ±3 36 ±18 37 ±3

In:

Here also the extent of the reaction with sodium methylate was greatest in the normal solution, which indicated a yield of 36 18 per cent. of the isomer containing negative chlorus. The yield of the same isomer determined by Holleman (fusion curve method) was 4200 per cent.

TABLE III.

III, Action of Sodium Meinflare on the Isomers Derived from m-Chlorotolueme.

(See Scheme 3.)

Mixture of Isomers (gram)	Normality NaOCH ₂ .	AgCl (gram)	Per Cent. Yield of I vomtrs \(\Lambda' + \Lambda''', \tau_1, \tau_2, \tau_5 + \tau_1, \tau_4 + \tau_5 \) 1. 3, 2-CH ₂ , Cl. NO ₂ C ₀ H ₂ ,	Average.
0 2257 0 2510 0 2672 0 2520 0 2570 0 3212 0 2809 0 2808	0'5 N 0'5 N N N 2 N 2 N 3 N 3 N	0°1652 0 1843 0 1987 0 1853 0°1898 0 2308 0 1923 0 1888	\$7:56 \$7:56 85:99 87:99 85:90 85:08 85:98 86:46	87 ,7 88:49 85:52 81 19

Again the extent of the reaction was greatest in the normal solution indicating a yeld of 88-40 per cent of the three isomers containing negative chlorine. Wibaut (fiactional crystallization and fusion curve methods) obtained a 100 per cent. yield of these isomers.

The yields of the isomers containing negative chlorine determined by physical methods and the maximum yields of the same isomers determined by the chemical methods just described are summarized in the following table:—

TABLE IV.

Scheme.	Isomers	Physical.	Chemical,	Percentage Ratio	١.
t 2 3	(A' + A'') (A') (A' + A" + A"')	62 20 (Wibaut) 42'00 (Holleman) 100'00 (Wibaui)	58 73 (Fry) 36'18 " 88 49 ",	94'42 86'74 88'49	

The extreme care with which Holleman and Wibaut conducted their physical determinations leads one to accept their results as practically final. Accordingly, in the last column above, the chemically determined yields are recorded as percentages of the physically determined yields. If each of the values in the last column was 100 per cent, then it might be asserted without question that the conception of the electronic tautomerism of the o-, o-, and m-chlorotoluenes given in schemes 1, 2, and 3, respectively, is completely substantiated by chemical methods. It may, nevertheless, be maintained that in so far as the values in the last column do approach the theoretical, so far, at least, do they substantiate both by physical and by chemical methods the electronic tautomerism of the chlorotoluenes. Moreover, it may be said that the electronic tautomeusm of o-, p-, and m-chlorotoluenes is the only hypothesis that has accounted for the replaceability of certain halogen atoms in some of the isomeric netrochlorotoluenes and the non-replaceability of certain halogen atoms in others. The data presented in this chapter substantiate the hypothesis

The fact that the chemically determined quantities of the isomers containing negative chlorine were less than the quantities determined by physical methods may have been due to two causes. First, the interaction with sodium methylate of one or more of these isomers in the mixtures may have been incomplete. This is not likely, since the results of the experiments with the analogous chloronitrobenzene gave results in perfect agreement with the theory. Furthermore, a survey of the quantitative data given in connection with schemes 1, 2, and 3 shows a maximum replacement in every reaction in which the unmormal solutions of sodium methylate were used. Concentrations below or above normal gave lower yields

In the second place, the low yields of the isomers containing negative chlorine may be due to the fact that the relative quantities of the isomers obtained on nitiation of the chlorotoluenes in Holleman's laboratories may not have been identical with the relative quantities of the same isomers obtained in this laboratory, even though the original directions given by Holleman and Wibaut were carried out precisely. The variations are most likely due to lack of standardization in certain details of manipulation, such as the rate of the addition of the nitric acid, methods of stirring, cooling, etc. While the temperature variations were kept within the prescribed limits, primarily to avoid the formation of poly-nitro-derivatives, it is quite likely that the local temperature

quantities of the isomers formed In fact, I have found that when nitiations were conducted separately, but under identical conditions, variations in the fusion points of the resulting mixtures were noted. This signifies corresponding variations in the relative quantities of the isomers in these mixtures

effects in the reaction mixture may have altered the relative

In conclusion, the variations between the chemically and the physically determined yields are not sufficient to invalidate the

principle of the electronic tautomerism of the chlorotoluenes, The conception affords a consistent interpretation of the observed

facts, and is substantiated in a great measure by the quantitative data offered in this chapter A one hundred per cent quantitative confirmation of the electronic tautomerism of the inixed dihalogen benzenes (notably para-chlorobromobenzene) is found in the experimental data presented in Chapter X Indeed, the agreement between experiment and hypothesis, definitely indicated in Chapters X, XI. and XII., establishes the principle of the electronic tautomerism of benzene derivatives This principle makes it possible to interpret the simultaneous formation of ortho-, para-, and meta- di-substituted, and poly-substituted derivatives of benzene.

yields of the ortho- and para-derivatives of $C_s^\dagger H_s$, \bar{X} will exceed the yield of the meta-derivative of $C_s^\dagger H_s$, \bar{X} . On the other hand, if electromer $C_s^\dagger H_s$, \bar{X} preponderates, then the yield of the meta-derivative will be greater than the combined yields of the ortho-and para-derivatives.

It has been assumed that the preponderance or increased concentration of one electromer over the other depends upon, or is a function of, the tendency of the given substituent to function

in one way rather than in another; thus, either $\hat{X}>\hat{X}$ or $\hat{X}>\hat{X}$. This tendency has been referred to, in the preceding chapter, as the polar stability of the substituent. Accordingly, the concentrations of the electromers are predetermined by the polar stabilities of the substituents present in the benzene nucleus.

It now becomes necessary to develop further this conception of polas stability and to inquire how and to what extent it predetermines the types of substitution, and, concomitantly, the quantitative yields of the substitution products To this end, the nitration of toluene, benzyl chloride, benzal chloride, and benzotrichloride, quantitatively investigated by Holleman, 88 affords significant data which are given in the following table:—

	C ₆ H ₅ X	Y.	Ortho.	Para.	(o + p.)	Meta.
2 3 4	C ₀ H ₈ . CH ₈ . C ₀ H ₈ . CH ₉ Ci C ₆ H ₆ . CHCl ₉ C ₆ H ₅ . CCl ₈ .	NO ₂ NO ₂ NO ₂ NO ₂	58 8 40.9 23.3 6 8	36·8 54·9 42·9 28·7	95 6 95 8 66*2 35 5	4'4 4'2 33'8 64'5

The first column embodies the formulæ of the mono-substituted detivatives, C_0H_0 . X, which are nitrated at or near o' Centigrade, under conditions that lead to the substitution of only one nitro group, Y, yielding in each of the nitration teactions (Nos. 1, 2, 3, and 4) varying quantities of ontho-para- and meta-nitrotluenes, C_0H_0 XY. The fifth column gives the combined yields of orthoand-para-derivatives, while the sixth column shows the corresponding yields of meta-nitrotoluene.

From the standpoint of the scheme for the nitration of a mono-substituted derivative, C_uH_uX , yielding simultaneously ortho-, para-, and meta-di-substituted derivatives, C_uH_uXY

substituent X to function negatively is greater in C_nH_1 , CH_2 , and in C_nH_2 , CH_2 Cl, than it is in either C_nH_2 , $CHCl_2$ or C_nH , CCl_2 . This tendency, or relative polar stability, is represented thus:

CH₂ or CH₂Cl>CHCl₂>CCl₂ Consequently, the combined yields of the ortho- and para-nitrobluenes, while practically the same in the nitration of C₂H₃, CH₄ and C₂H₃, CH₄Cl, are found to decrease when C₄H₃, CHCl₃ and C₄H₃, CCl₄ are initrated In other words, the concentrations of the electromers of the type

 C_0^{-1} , X in the successive nitrations 1, 2, 3, and 4 (preceding table) are relatively as follows —

$$C_{c}^{+}H_{s}$$
. CH_{2s}^{-} or $C_{c}^{+}H_{s}$. $CH_{2}^{-}CI > C_{c}^{+}H_{s}$. $CHCl_{s} > C_{c}^{+}H_{s}$. CCl_{s} .

On the other hand, it may be noted that the tendency for the substituent CCl₄ to function positively is greater than that of either CHCl₂ or CH₂Cl and CH₂. These relative polar stabilities

are indicated thus: $\overrightarrow{CCl_s} > \overrightarrow{CHCl_s} > \overrightarrow{CH_s} \overrightarrow{Cl}$ or $\overrightarrow{CH_s}$. Consequently, the yields of meta-nitrotoluene increase in the successive nitrations as noted in the preceding table. In equivalent terms, the concentrations, or reactivities, of the electromers of the

opposite type, $C_4H_4^+$ X, in the successive nitrations (I, 2, 3, and 4) are related as follows .—

$$C_0\vec{H}_5\,.\,C\overset{+}{H}_5\,\,or\,\,C_0\vec{H}_5\,.\,C\overset{+}{H}_5Cl>C_0\vec{H}_5\,\,\,C\overset{+}{H}Cl_1>C_0\vec{H}_5\,.\,C\overset{+}{Cl},$$

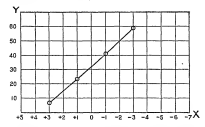
The conception of polar stability as heretofore applied to substituents or radicals does not lend itself to mathematical definition, but there may be assigned to a given substituent a polar number which may be defined as the algebraical sum of the positive and negative valences of that atom of the substituent or radical which is united to the benzene nucleus. Since a given substituent may function either positively or negatively, there shall not be included in this polan number the positive or the negative valence which unites the radical to the nucleus According to this definition, the polar numbers of the radicals CH₃. CH₃Cl, CHCl₂, and CCl₃ appear from their electronic formula—

to be, respectively, - 3, - 1, + 1, and + 3.

A remarkable relationship is found to exist between these polar numbers and the quantitative yields of the ortho-nitio-derivatives. In the following table, the first column embodies the formulæ of the benzene derivatives, C_cH_sX , subjected to nitation. The second column gives the polar number of the substituent X_i , and the third column contains the corresponding percentage yields of the ortho-nitro-derivative:—

C _d II ₅ .X	Polar Number of Substituent X.	Ortho-miro- derivative.
C,H ₅ .CH ₃	- 3	58°8
C,H ₅ .CHCl ₂	- 1	40°9
C,H ₅ .CHCl ₂	+ 1	23°3
C,H ₆ .CCl ₃	+ 3	6°8

If, in the system of rectangular co-ordinates, the polar numbers of the substituent (X) are indicated on the X axis and the corresponding percentage yields of the ortho-nitro-derivatives are indicated on the Y axis.



it is quite evident that the quantitative yields of the ortho-nitrosubstitution products of toluene, benzyl chloride, benzal chloride,

numbers of the respective mono-substituents, CII, CH,CI, C-HCI, and CCI₄ present in the nucleus. In the same manner (rectangular co-ordinates) it follows that the combined yields of the ortho- and para-nitrotoliuenes on the one hand, or the yields of the meta-nutrotoliuene on the other hand, are also linear functions of the polar numbers of the substituents CH₂CI, CHCI, and CCI.

The existence of the foregoing relationships warrants a search for other quantitative relationships. To this end, the only quantitative data available are chiefly those of Holleman and his co-workers, who have carefully determined the yields (in per cent.) of the products of nitration of various mono-substituted derivatives of benzene. The nitration reactions were conducted so as to yield only mono-nitro-derivatives, orthor, metar, and para. All the available data are embodied in the following table:—

Substituent (%) in Nucleus	lf. Polar Number	III. Ortho	IV Para.	v (+ + †)	NI Meta.
CH ₄ CH ₂ Cl. le Cl. Br CHCl ₂ CCl ₄ CO ₂ C,H ₅ CO ₂ CH ₉ . CO ₂ CH ₉ .	- 3 - 1 0 0 0 + 1 + 3 + 3 + 3 + 4	56 0 40 9 12 1 30 1 37 6 41 1 23 3 6 8 28 3 21 0 18 5 6 8	40°9 54 9 87 6 69 9 62 4 58°7 42 9 28°7 3°3 5°8 1°2 0°0	96 9. 95 % 100 0 100 0 100 0 100 0 100 0 66 2 35 5 31 6 26 8 19 7 6 8	3.1 4.2 0.0 0.0 0.0 0.0 0.0 33.8 04.5 65.4 73.2 80.3 93.2

Column 1 embodies the formulae of the substituents (X) of the mono-substituted derivatives C_0H_0X . The polar numbers of these substituents are indicated in column II. Columns III., IV., and VI. embody the percentage yields of the ortho-paus, and meta-nutro-compounds, respectively, of general formula C_0H_0 , X. NO_2 . In column V the sum of the percentage yields of the ortho- and para-isomers is given. The data of the table have been tabulated in the orde of the increasing values of the polar numbers of the substituents, i.e., from -3 to +4. This arrangement reveals some striking regularities which may be summarized as follows -0.

- (1) When the polar number of a given substituent is zero or less than zero the combined yields of the ortho- and paa:nutic-compounds range between innety and one hundred per cent. In other words, the ortho-para type of substitution and percentage yield of products predominate practically to the exclusion of the meta type of substitution and yield when the polar number of the substituent in the nucleus is zero or less than zero.
- (a) As the polar number assumes a positive sign and increases in value, the corresponding ortho-para percentage yield decreases while the meta yield increases. Thus when the polan number is + 4 the meta type of substitution and yield predominate almost to the exclusion of the ortho-para type and yield.
- (3) From (1) and (2) it appears that the combined yields of the ortho- and para-nitro-substitution products and the yield of the meta-nitro-substitution product are predetermined, within certain limits, by the polar number of the substituent X in C_eH_eX. Therefore, if the yields of the ortho- and para-derivatives are dependent upon the speed of interaction of the electromer

of the type $C_{\mathbf{b}}^{c}H_{\mathbf{b}}$. X and the yield of the meta-derivative depends upon the speed of the interaction of the electromer of the type

- C_nH_n , \dot{X} in the tautomeric system C_nH_n , $\dot{X} \Leftrightarrow C_nH_n$, \dot{X} , then it follows that the speed of the interaction of these electronners with the substituting reagent is a function of the polar number of the substituting \dot{X} in C_nH_n . \dot{X} . Furthermore, if the speeds of reaction of the electronners are dependent upon their concentration, then these concentrations are also a function of the polar number of the substituents.
- (4) Finally, it should be noted that, in terms of positive and negative valences, the polar number of a substituent is in eality an index of the degree or state of exidation of that atom of the substituent which is directly united to the benzene nucleus. This is inhamony with the principle that the development of positive valences (i.e., loss of electrons) concesponds to oxidation while the development of negative valences (i.e., gain of electrons) corresponds to reduction. In other words the problem of benzene substitution appears to be resolving itself into a study of the states of oxidation or reduction of the atoms of the substituents, i.e., that atom of the substituent which is in direct combination with a carbon atom of the benzene nucleus. It is

significantly interesting to recall, inducetly in this connection, that the empirical rule of Brown and Gibson was based upon the susceptibility to oxidation of certain compounds. The significance of this feature was interpreted in the further analysis of the Brown and Gibson rule (Section B. Charlet IX.).

The present chapter has indicated the existence of a relationship between the polar number of the substituent X in $C_0H_0X_0$ and the combined quantitative yields of the ortho- and paramono-nitro-substitution products on the one hand, and the quantitative yield of the meta-intro-derivative on the other. There yet remains to be developed a relationship between the variable quantitative yields of the ortho- and the para-mono-nitro-substitution products, and the respective electronic formulæ of these products. Perhaps a solution to this problem may also be lound in a further development of the conception of the polar numbers of the substituents.

Now in order that formic acid, in which three of the carbon valences are positive, and one is negative, may yield caibon dioxide, in which the four valences of carbon are positive, the one negative valence of carbon in formic acid must be oxidized to a positive valence. This change takes place when formic acid is catalytically or photochemically decomposed, yielding carbon dioxide and hydrogen according to the following electronic equation —

Here also the photochemical change is an intramolecular oxidation-reduction reaction, the electromer $H^+ - CO_2H$ is transformed to the electromer $H^- + CO_2H$. Then decomposition

formed to the electromer H——CO₂H. Then decomposition into carbon dioxide and hydrogen is possible and takes place. In this chapter some fundamental distinctions between nucleus

and side-chain substitution have been developed and interpreted in terms of positive and negative valences. A common occurrence, however, should not be overlooked, namely, that in most substitution reactions it is practically impossible so to limit and regulate the conditions that the sole reaction taking place will be either nucleus substitution or side-chain substitution. In other words, the formation of ortho- and para-halogen substituted toluenes, in which the halogen atoms are positive, and the introduction of negative halogen into the side-chain, may pioceed as simultaneous independent chemical changes. As previously noted the former process does not involve oxidation and reduction while the latter does. These facts are readily correlated

with another general property characteristic of benzene on the one hand, and characteristic of aliphate hydrocarbons (comparable to side-chains) on the other hand. For conversity ordation, while the aliphatic hydrocarbons and the side-thains are generally more susceptible to audation.

Finally, it is not impossible to conceive that, under certain conditions, which are not as yet determined and interpreted, a positive hydrogen atom of the benzene nucleus may be replaced by a negative halogen atom (or vice versi), which is characteristic of side-chain substitution, involving oxidation and reduction. Sucha condition, however, does not invalidate the interpretations presented in this paper, which are correlated with the experimental facts, the conditions of substitution, and the chemical properties of ortho- and para-halogen substituted toluenes, and the side-chain substitution products.

In concluding this chapter, the relations between the electionic interpretations and the substitution hypotheses of Bruner, Bancroft, and Holleman should be reconsidered briefly

Bruner's idea that nucleus substitution is due to halogen atoms, while side-chain substitution is due to molecules, thus,

$$C_0H_1CH_1 + 2Br \Rightarrow BrC_0H_1CH_1 + HBr,$$

 $C_0H_1CH_1 + Br \Rightarrow C_0H_1CH_1Br : HBt,$

fails to take into account the fact that nucleus substitution (former equation) does not involve oxidation and reduction, while side-chain substitution (latter equation) does.

Bancont's assumption that nucleus substitution is due to negative halogen atoms is not in agreement with the evidence, both theoretical and eveperimental, that the othos and parasubstituted halogen atoms of toluene are positive. His other assumption that the halogen substituted in the side-chain is positive is also contradicted by the fact that the halogen atoms in benyl chloride, bencal chloride, and benzotrichloride (or bromides) are negative, since they are readily exchanged (without oxidation or reduction) for negative hydroxyl. It is true, however, that positive halogen atoms are the active substituting azents in side-chain substitution, but only by virtue of their oxidizing action and consequent reduction to negative halogen

thèse, suivant laquelle le noyau est attaqué pai des molécules de $\overline{HBr_m}$, tandis que la chaine latérale l'est par les molécules de brome, a quelque viaisemblance; cependant un nombre de difficultés doivent encore être levées, avant qu'elle puisses servir à expliquer les phénomènes obseivés". The very plausible interpretation of the nucleus bromination of toluene though the action of the intermediately formed $\overline{HBr_m}$ may be readily correlated with the interpretation of nucleus substitution, and the action of halogen carriers as presented in this chapter, provided action of halogen carriers as presented in this chapter, provided



chain substitution is subject to the criticism that it also fails to recognize the side-chain substitution process as an oxidation and reduction phenomenon depending upon the reaction,

X + 2⊕, previously considered.

CHAPTER XVI.

"FREE RADICALS" THEIR EXISTENCE AND PROPERTIES

RADICALS are conceived to be groups of atoms which function as single atoms in chemical relations and, accordingly, preserve their composition or constitution throughout chemical changes. The history of Chemistry affords many instances of attempts to isolate radicals, i.e., to pieque "free adicals". As early as 1831 Liebiq, "suggested the possibility of isolating, and seriously attempted to piequate, free radicals. In 1830 Lowig "claimed to have effected the separation of the radical ethyl, Cyfle. An instorical review of these various attempts, chiefly failures, would constitute an interesting chapter, the culmination of which, however, is found in the recent work of Gomberg, who, undoubtedly, has isolated free radicals, notably in the form of the triarr/inethyls.

The triarylinethyls are regarded by many to be compounds of tervalent carbon Schlenck's discoveries, by means of molecular weight determmations, that some of the triarylinethyls are largely mono-moleculal (for instance, tri-bi-phenylinethyl was found to be entirely mono-molecular) leads, as Gombeig has emphasized, to but one inference, namely, that these compounds exist as fice radicals, i.e., they are compounds of the tervalent carbon atom of the general formula (R) = C

The present chapter offers an interpretation of the existence and properties of free radicals in terms of the electronic conception of positive and negative valences which reveals the existence of many significant relationships that are apparent in electronic formulae but that cannot be found in ordinary structural formulae. The principles developed in Chapter IV, Electronic Amphotosism, and especially those of Chapter V, The Nascent State, are both essential and fundamental to an appreciation of the interpretations presented in this chapter.

Particular attention should be redirected to the fact that all

the chemical actions classed as "nascent" are of a reduction or an oxidation type, and that the so-called "nascent state" is conceived to be an initiable condition of a substance which manifests (anter certain conditions) an adaptibility and a tendency to lose electrons, or gain electrons, and thereby revert to a more stable condition. If the substance (ion, atom, or molecule) lose negative electrons, it acts as a reducing agent. If it combines with negative electrons, it acts as an ovidining acent.

From the point of view embodied in the above definition, the present chapter will endeavour to show that free radials such as the trarylmethyls, are comparable in many respects with the electroally neutral atomic state, and that their instability, or reactivity, and general chemical properties may be interpreted by their development of positive or of negative valences through the loss or gain, respectively, of negative electrons. In other words, the triarylmethyls may act either as reducing or as oxidizing agents. Furthermore, in explaining some of the reariangement reactions peculian to triphenylmethyl, the electronic formula of benzene will be shown to play a significant part.

Consider, first, the reaction for the preparation of triphenyl-methyl. Gomberg ¹⁰⁰ submitted triphenylchloromethane to the action of metals, notably silver, with the full expectation that the reaction would proceed normally and give rise to hexaphenyl-ethane:—

He obtained "a hydrocarbon possessing the requisite composition (C = 93'8 per cent, H = 6'2 per cent,)... But the unusual instability of this substance, its proneness to enter into the most varied chemical reactions, and above all, its striking unsaturated character, precluded the natural inference that the hydrocarbon at hand was actually hexaphenylethane. The opinion was expressed that here was an instance of a compound with one atom of carbon in the trivalent state, i.e., $(C_0H_1)_C$, triphenylmethyl, a free radical." Now it may be shown that the electronic interpretation of the action of silver upon triphenylchloromethane also substantiates the assumption of the existence of the free radical as a compound of trivalent carrior.

Triphenylchloromethane presumably possesses the abbievi-

ated electronic formula (C₆H_c)_cC —Cl. Its aqueous solution is a conductor of the electric current depending upon ionic dissociation—

Furthermore, its hydrolysis conforms to the equation

$$(C_aH_g)_a\overset{+}{C}$$
, $\overset{-}{C}I$ + $\overset{+}{K}$, $\overset{-}{O}II$ \rightarrow $(C_aH_g)_a\overset{+}{C}$ $\overset{-}{O}H$ + $\overset{+}{K}$, $\overset{-}{C}I$,

Metallic silver (with zero valence) in order to combine with negative chlorine, must itself develop a positive valence, i.e., it must become positively univalent through the loss of an electron

 $Ag \rightarrow \bigoplus + Ag$ Accordingly, the electronic scheme for the action of silver upon triphenylmethylchloride is represented as follows:—

$$(C_0H_0)_tC$$
 $\stackrel{\leftarrow}{-}CI \Rightarrow (C_0H_0)_tC$
 $\stackrel{+}{C}I \stackrel{\leftarrow}{+}CI$
 $Ag \Rightarrow \bigoplus_{\substack{\downarrow \\ (C_0H_0)_tC}} Ag \stackrel{\downarrow}{+}CI$

It should be observed that the methyl carbon atom in the triphenylmethyl ion is quadrivalent, but the free positive valence of this ion is obliterated by the acquisition of the negative electron from the metallic silver. In other words, silver has reduced the quadrivalent methyl carbon atom to the tervalent state, yielding triphenylmethyl.

The possible reaction of two electrically neutral molecules of triphenylmethyl to form hexaphenylethane is exactly parallel to the reaction of two electrically neutral hydrogen atoms to form molecular hydrogen. The reaction depends upon one of the atoms, or free radicals, losing an electron which is acquired by the other atom, or free radical, respectively. Combination of the resulting oppositely charged atoms or radicals occurs smultaneously as indicated in the following scheme:—

³ See the equally applicable "quinocarbonium formula" in a later section of this chapter, p. 161.

The following properties of triphenylmethyl will be considered: (A) oxidation; (B) addition of iodine; (C) addition of hydrogen; (D) leaction with nitrogen dioxide (or tetroxide); (E) addition of xylene, (F) action of acids; (G) action of light; (H) electrical conductivity of triphenylmethyl when dissolved in liquid sulphur dioxide, based upon the analogous conductivity of sodium dissolved in liquid ammonia

Gomberg has frequently called attention to the fact that triphenylmethyl behaves in some reactions as if it were an element of basic nature. This is particularly so in its reactions with oxygen and with iodine. This behaviour, from the electronic standpoint, may be attributed to the tendency of the tervalent carbon atom to become quadrivalent, i.e., to develop a positive valence through the loss of an electron in just the same way that metallic silver or sodium develops a positive valence:—

$$Na \rightarrow \stackrel{+}{Na} + \bigcirc$$
.
 $(C_0H_0)_3C \rightarrow (C_0H_0)_3\stackrel{+}{C} + \bigcirc$.

The striking similarity of triphenylmethyl to sodium is shown in the reactions of each with (A) oxygen, and (B) iodine.

A. Oxidation.

On exposure of a concentrated solution of triphenylmethyl in benzene (10 per cent.) to air, a colourless fauly stable peroxide, {(C₈H₃)₂C₃, is formed Sodium and oxygen combine at temperatures below 180° yielding a mixture of sodium oxide and peroxide As the temperature is raised the quantity of peroxide increases At 300° sodium peroxide is the principal product. Now metallic sodium possesses neither positive nor negative valences, but in its oxide and peroxide sodium is presumably positively univalent, since these oxides on hydiolysis

yield sodium hydroxide, Na. OH. Accordingly, the complete electronic scheme for the formation of sodium peroxide may be remessated as follows.—

The parenthetical part of the scheme involves the reduction of a positive valence of oxygen to a negative valence by the action of the electrons from sodium. The sodium atoms, thus becoming positive, unite with the negative valences of the oxygen atoms.

In a perfectly analogous manner triphenylmethyl unites with oxygen, thus:--

$$2(C_1H_3),C$$
 + O_4 $\sim (C_4H_3),C$ O_5 $\sim (C_4H_3),C$ O_6 O_7 O_8 O_8

B. Addition of Iodine.

The combination of either sodium of triphenylmethyl with iodine is also an oxidation-reduction tenction. Gomberg ¹⁵⁸ has shown that triphenylmethyl may be titrated with a standard solution of iodine. Triphenylmethyliodide is formed. The iodine in this compound is negative since it is replaced by negative hydroxyl on hydrolysis. The iodine atom in sodium iodilide is also negative since the aqueous solutions yield the ions

Na and I Accordingly, the electronic schemes for the combinations of sodium and of triphenylmethyl with iodine (i.e., then oxidation by means of iodine) are represented as follows.

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In the parenthetical part of the scheme the elections from sodiumreduce I to I. In the following scheme the elections from tri-

phenylmethyl accomplish a like change :-

In the preceding reactions (1) and (2), the tervalent methyl carbon atom of triphenylmethyl functioned as a reducing agent by losing an electron, and thereby becoming quadrivalent through the development of a positive valence. On the other hand, when triphenylmethyl combines with hydrogen, the methyl caubon atom becomes quadrivalent through the acquisition of an electron, i.e., the development of a negative valence. In this respect, triphenylmethyl may be said to resemble certain non-methy

C. Addition of Hydrogen.

Previously it has been noted that only negative hydrogen (H)

or neutral hydrogen (H) acts as a reducing agent: $H \rightarrow H + \Theta$;

 $H \rightarrow \dot{H} + \bigcirc$. Positive hydrogen does not act as a reducing agent because the further loss of an electron would render

hydrogen bivalent, an unknown condition: $H \rightarrow H + \bigoplus$ Hence, the interaction of the electrically neutral triphenylmethyl and neutral atomic hydrogen is perfectly analogous to the union of chloime and hydrogen atoms —

$$H + CI \rightarrow HCI$$
 $\downarrow + H + O + CI \rightarrow H \stackrel{+-}{\longrightarrow} CI$

Atomic hydrogen reduces atomic chlorine to negative chlorine, or, in equivalent terms, atomic chlorine oxidizes atomic hydrogen to positive hydrogen. In the following scheme, atomic hydrogen reduces the tervalent methyl carbon atom, which, acquiring a negative valence, becomes quadrivalent in triphenylmethane:—

D. Reaction with Nitrogen Dioxide (or Tetroxide).

Schlenck and Mair 182 found that nitrogen dioxide on interaction with triphenylmethyl yielded simultaneously two products as indicated in the equation—

$$2(C_0H_1)_3C \ + \ 2NO_2 \ (\text{or} \ N_3O_4) \ \Rightarrow \ (C_0H_3)_3C \ . \ NO_0 \ + \ (C_0H_3)_3C \ . \ NO_0 \ (\text{triphenyl-ntrone-thane})$$

In the electronic interpretation of this interesting reaction, the former compound must be regarded as a derivative of nitric acid, $HO = ^+NO_2$, the latter, a derivative of nitrous acid, $H = ^+NO_0$, or $H = ^+O = ^+NO$. The key to the interpretation is found in the fact that introgen tetoxide is hydrolyzed by water yielding nitric and introus acids. The following scheme takes into account the tautomeric formulae of N_2O_4 , to which correspond the tautomeric forms of nitrous acid :-

$$N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} + \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0 - \frac{1}{N_{0}} = 0$
 $N_{0}Q_{0} \approx 0$

Each of the five valences of the introgen atom in intric acid is positive, but in introus acid four are positive, and one is negative. Hence, in the tautomer H—O—N=O, nitrogen is tet valent since one positive and one negative valence are polarized. Note also that one intro-radical functions positively, while the other (either NO₂ or ONO) functions negatively. Therefore, when one molecule of triphenylmethyl loses an electron and

runctions positively, another molecule of triphenylmethyl acquires that electron and functions negatively. Consequent union of negative triphenylmethyl with the positive intro-radical yields triphenylnitromethane, while union of positive triphenylmethyl with the negative nitro-andical yields triphenylmethylmithethyl with the negative nitro-andical yields triphenylmethylnitrite. These changes are indicated as follows—

$$a(C_{k}H_{k})_{k}C \rightarrow (C_{k}H_{k})_{k}C$$
 $+ (C_{k}H_{k})_{k}C$
 $+ (C_{k}H_{k})_{k}C$
 $+ (C_{k}H_{k})_{k}C$
 $+ NO_{k} + NO_{k} \text{ or } ONO_{k}$
 $+ NO_{k} \text{ or } ONO_{k}$
 $+ NO_{k} \text{ or } (C_{k}H_{k})_{k}C$
 $+ NO_{k} \text{ or } (C_{k}H_{k})_{k}C$
 $+ ONO_{k} \text{ or } (C_{k}H_{k})_{k}C$

It should also be noted that, depending upon conditions, the dissociation of nitrogen tetroxide may proceed either ionically or molecularly. In aqueous solution, the dissociation is pre-

sumably ionic, $N_2O_4 = (NO_3)^{--} - (NO_2) \Rightarrow NO_2 + NO_2$, since hydiolysis yields nitric and nitious acids as indicated above. But when nitrogen tetioxide is heated to 140^5 , its density corresponds to the dioxide formula. Is it not possible that NO_9 , in which nitrogen is quadrivalent, exists as a free indical just as $(C_4H_5)_4C$ is a free radical in which arbon is tervalent? O_7 , to not the dissociation of associated triphenylmethyl (hexaphenylethane) exactly parallel to the dissociation of associated nitrogen dioxide (nitrogen tetroxide)? This is quite evident from the following equilibility a^{--}

$$\begin{split} (C_gH_g)_gC \cdot C(C_gH_g)_3 &= (C_gH_g)_3 \frac{+-}{-}C(C_gH_g)_3 & \rightleftharpoons (C_gH_g)_3C \\ & N_gO_4 &= (NO_2) \frac{+-}{-}(NO_2) & \rightleftharpoons NO_2 \\ &+ NO_2. \end{split}$$

In each of the above non-ionic dissociations, the positive radical becomes neutral, or is reduced to a free radical, through the acquisition of an electron from the negative radical which is thereby also rendered neutral, i.e., it is oxidized to a free radical.

These points of view may be summarized as follows In notic dissociation $(A \xrightarrow{+} B \Rightarrow A + B)$ the atoms or radicals maintain their respective charges. In molecular dissociation

 $(A - B \rightleftharpoons A + B)$ the atoms or radicals become electrically neuthal through the transference of electrons from the negative radical to the positive radical. Thus, fice radicals are formed, and their activity is a factor of their further tendency either to gain or lose electrons. In electronic tautomerism ¹⁸².

a further transference of electrons reverses the polarity of the constituent atoms or radicals, and in this change the intermediate, electrically neutral atoms or molecules (A and B) are escentially five radicals. Three younts of view wive to emphasize the definition of a five radical as an electrically ratified above molecule which is capable of developing either positive or negative volumes depending upon extarn conditions which effect the loss or gain of elations.

In the following interpretations of (E) the addition of xylene, (F) the action of acids, and (G) the action of light, the electronic formula of benzine plays an essential part.

E. Addition of Xylene.

Wieland and Muller wit found that the heating together of triphenylmethyl and xylene yielded dimethyltetraphenylmethane.

 $\neg (C_2H_3) C + C_4H_4(CH_3) = (C_4H_3) C, C_4H_3 \neg (CH_3) + (C_4H_4) CH$. They interpreted the reaction as proceeding in two stages: first, the union of two molecules of triphenylmethyl with one molecule of xylene; second, the elimination of one molecule of triphenylmethyl with a hydrogen atom of the benzene nucleus. These reactions are indicated in the following scheme which employs centric formula: —

$$\begin{array}{c} CH_1 \\ H - \\ H \end{array} \begin{array}{c} CH_2 \\ + 2 (C_4 H_3)_2 C \\ + \\ CH_2 \end{array} \begin{array}{c} CH_2 \\ + \\ CC_4 H_3)_2 C \\ + \\ CC_5 H_3 \\ + \\ CC_$$

Every detail of this scheme may be correlated perfectly with the electronic formula of benzene, and with the previously described mechanism of substitution involving the centric valences of the benzene nucleus. It has been shown that, as a general rule, when hydrogen atoms or substituents in positions 1, 3, and 5 are negative, hydrogen atoms or substituents in positions 2, 4, and 6 are positive; and vice versa Furthermore, the polarity of a centric valence of a given carbon atom of the benzene nucleus was shown to be opposite in sign to the polarity of the hydrogen atom or substituent united to that carbon atom. Now the capacity of the methyl carbon atom of triphenylmethyl to develop either a positive or a negative valence has been demonstrated in preceding paragraphs. Accordingly, the addition of two molecules of tuphenylmethyl to two of the centric valences of the nucleus, depends upon one molecule of triphenylmethyl functioning positively while the other molecule functions negatively The electronic interpretation of the interaction of tirphenylmethyl and xylene is embodied in the following scheme -

$$\begin{array}{c} H \stackrel{+}{\longrightarrow} H \stackrel{+}{\longrightarrow} H \\ CH^{2} & C(C^{1}H^{2})^{2} & H \stackrel{+}{\longrightarrow} H \\ CC^{1}G^{1}H^{2} & C(C^{1}H^{2})^{2} & C(C^{1}H^{2})^{2} \\ CH^{2} & CH^{2} & CH^{2} & C(C^{1}H^{2})^{2} \\ CH^{2} & CH^{2} & CH^{2} & C(C^{1}H^{2})^{2} \\ CH^{2} & CH^{2} & CH^{2} & CH^{2} \\ CH^{2} & CH^{2} \\ CH^{2} & CH^{2} & CH^{2} \\ CH^{2} \\ CH^{2} \\ CH^{2} & CH^{2} \\ CH^{2} \\ CH^{2} \\ CH^{2} \\ CH^{2}$$

From the above it is evident that the interaction of triphenylmethyl and xylene is not, strictly speaking, an addition reaction, but rather a substitution reaction conforming to, and thereby substantiating, not only the mechanism of substitution presented in previous chapters, but also the capacity of triphenylmethyl to function as a positive ion and as a negative ion.

In considering (IF) the action of acids, and (G) the action of light upon triphenylmethyl, the electronic formulæ of the benzene nuclei of triphenylmethyl are involved. The phenyl radical is commonly spoken of as being negative in character, but if two phenyl radicals are united directly, as in diphenyl, the electronic conception of valence regards one phenyl as negative while the

othen is positive $(C_6H_3)^{1}$ — (C_cH_3) . Now triphenylmethane is generally made, either from benzal chloude, or from chloroform, through replacement of chloune by phenyl. Hydrolysis of either of these halogen compounds by potassium hydroxide effects the replacement of chloune by negative hydroxyl. From this standpoint, the chlorine atoms function negatively, and their replacement by phenyl radicals would indicate that the latter are also negative in triphenylmethane, and in triphenylmethyl. Hence the electronic formula for triphenylmethyl, in which the valences of only one of the phenyl groups need be indicated, is as follows.

It should be observed that the hydrogen atoms in the orthoand para-positions are negative. The tendency for negative hydrogen to lose an electron and become neutral $(H \to \bigcirc + H)$ and the further tendency for neutral hydrogen to lose an electron,

and become positive $(H \rightarrow \bigcirc + \overset{\circ}{H})$ has been considered previously in connection with various reactions and interpretations.

Therefore, it may naturally be assumed that the negative hydrogen atoms of the benzene nucleus are possibly less stable, in the above sense, than the positive hydrogen atoms. Moreover, if we consider, for example, the negative para-hydrogen atom in the above formula of triphenylmethyl, what would be the immediate consequences of its becoming neutral through the loss of an election? The positive valence of the para-carbon atom would be obliterated (neutralized or polarized) by the negative electron from the para-hydrogen atom. In other words, the para-carbon atom would be reduced to the tervalent state, and the resulting neutral hydrogen atom would be free or dissociated But its further tendency to lose another electron and thereby become positive might naturally lead to its immediate attachment to the unsaturated trivalent methyl-carbon atom which would develop a negative valence through the acquisition of the electron from the neutral hydrogen atom, and thereby hold in combination the resulting positive hydrogen atom. These changes (intramolecular rearrangements) are illustrated in the following scheme :-

$$\begin{aligned} & Formula \ I \\ & H \\ & H$$

This scheme, based upon the tendency of negative hydrogen to function positively, affords both an explanation and a mechanism of the wandering of the para-hydrogen atom to the

methyl carbon atom, and also shows how the para-carbon atom in formula I for triphenylmethyl is rendered tervalent in the tautomeric modification of formula II. In other words, the methyl carbon atom of tautomer L is trivalent, the para-carbon atom of tautomer 11, is fervalent. The existence of tautomer 11. readily lends itself to, and is the essential feature of, the followme interpretation (F) of the action of acids upon triphenylmethyl,

F Action of Acids

Concerning the action of acids upon triphenylmethyl, Gomberg states that "Hydrochloric acid, and undoubtedly other acids, exert a peculiar catalytic influence upon triphenylmethyl. The musaturated, unstable hydrocarbon is thus converted in the presence of small quantities of acids, into an isomeric stable hydrocarbon. This hydrocarbon was first prepared by Ullmann and Borsum, directly from triphenylchloromethane, and later by the writer from triphenylmethyl as mentioned above. It was first taken for the true stable hexaphenylethane. But Tschitschibabin definitely proved its constitution to be that of A-benzylhydryltetraphenylmethane. The rearrangement takes place thus .--

$$_2(C_oH_s)_1C \rightarrow (C_oH_s)_2CH - \langle --- \rangle - C(C_gH_s)_1$$
,"

This reaction is readily interpreted by the existence of tautomer II. (see preceding scheme) in which the para-carbon atom of one of the phenyl radicals is in the tervalent state The union of this carbon atom of tautomer II, with the tervalent methyl carbon atom of tautomer I would naturally take place because the para-carbon atom originally possessed a positive valence, and, therefore, tends to lose an electron, and thus again develop a positive valence. In turn, this electron would be taken up by the unstable tervalent methyl-carbon atom of tautomer I. Combination of the resulting oppositely charged tautomers would then yield parabenzhydryltetraphenylmethane according to the following scheme -

G. Action of Light.

Concerning the action of light, Gomberg has stated that "Solutions of triphenylmethyl in benzene are quite susceptible to light, especially direct similght. Like most photochemical processes of this type, the reaction in this instance also is auto-oxidation and reduction. Schimdlin showed that the following reaction takes place:—

$$3(C_0H_0)_0C \implies 2(C_0H_0)_0CH + C_0H_1 C - C_0H_0$$

Schmidlin, 105 employing bi-molecular formulæ, has presented the following scheme for this reaction .—

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

and he states that "das leicht dissoziierbare Hexaphenyl-athan wild durch das Triphenylmethyl zum schwer dissoziierbaren, energieaimen Di-biphenylendiphenylathan reduzieit"

Now the electronic formula for triphenylmethyl readily lends itself to an interpretation of this hitherto unexplained reaction. First, it should be observed that not only the hydrogen atoms in the paia-positions to the attachment of the methyl carbon atom, but also the ortho-hydrogen atoms, are negative. On the other hand, the methyl hydrogen atoms, are negative. On the other hand, the methyl hydrogen atom of triphenylmethane (one of the reaction products) is positive. Therefore, the auto-oxidation and reduction effected by high trivolves the transition of negative hydrogen atoms from the ortho-positions to the unsaturated methyl carbon atom of triphenylmethane. Here again the relatively unstable negative ortho-hydrogen atoms first become

neutral, $H \Rightarrow H + \Theta$, and the negative electron obliterates the costive valence of the ortho-carbon atom thus making it tervalent. The neutral or free hydrogen atom becomes positive

through the loss of an electron, $H \rightarrow H + \Theta$, which renders the unsaturated methyl carbon atom of triphenylmethyl negative Union of negative triphenylmethyl with positive hydrogen then yields triphenylmethane, and the union of the ortho-tervalent carbon atoms, by the transfer of an electron from one to the other, completes the formation of biphenylenedphenylethane. Employing the briefer unimolecular formula, the electronic interpretation is embodied in the following schemes (f) and (2). The first shows the effect of the conversion of the othen-negative hydrogen atoms to the free or neutral state, and the union of the resultant ortho-tervalent carbon atoms, which reaction is an intrimolecular arisation and robustion. The second shows the educing action of the neutral hydrogen upon triphenylmethyl vielding triphenylmethyl vielding triphenylmethyl

Thus the action of light in this particular instance involves

the change, $H \Rightarrow \dot{H} + 2\Theta$. In this connection it may be of interest to recall that in the preceding chapter the action of light in effecting nucleus substitution in the side-chain was shown to involve the conversion of positive halogen to negative halogen, $\dot{C} \Rightarrow Cl + 2\Theta$. These processes are analogous in that each

H. Electrical Conductivity.

involves auto-oxidation and reduction

Gomberg found that the triarylmethylhalides behaved truly like salts in that they manifest molecular conductivities which increase with the dilution. He states: "It was thus definitely established that there are 'carbonium' salts in the true sense of the definition applied to salts. The conclusion was drawn that the basic properties, by virtue of which the salt formation occurs, reside in the central methyl carbon atom," This conclusion is still held by some, but Gombeig later gave up this interpretation, presenting evidence which favoured the guino. hypothesis Accordingly, he states that "since the manifestation of colour and the salt-like properties of these substances are simultaneous, the conclusion seems warranted that both these phenomena are the results of one and the same cause, i.e., tautomerization to the quinod state. Consequently, these salts were named quinocarbonium salts corresponding to the quinocarbonium bases.

$$(C_aH_a)_iC$$
. OH $(C_aH_a)_2 = C = \bigcirc H$
(Carbinol, pseudo-base) (Quinocarbonium base)

"For obvious reasons, the basicity was now assumed to lie in the quinone nucleus, in the C* and not in the central carbon atom as had been originally supposed."

From these two points of view, triphenylmethylchloride (or other halides) may be represented electronically as carbonium or quinocarbonium salts by the respective formulæ—

$$(C_0H_5)_0C \stackrel{+}{-} X$$
 $(C_0H_5)_2 = C = X$

Since hydrolysis with potassium hydroxide yields potassium halide and the base, it follows that the triphenylmethyl ion (either formula) is positive. This brings us to consider the "unexpected discovery that triphenylmethyl itself behaves like an electrolyte when dissolved in liquid sulphur dioxide," "ion and concerning which Gomberg has raised the question in these words: "How is this stange phenomenon, a hydrocarbon behaving like an electrolyte, to be accounted for? Sevenal explanations have been offered, but none of these is entirely satisfactory."

Now since triphenylmethyl has been shown to act similarly to metallic sodium in several reactions (notably Section A and B of this chapter) for which the electronic interpretations were

based upon the parallel tendencies—Na \rightarrow Na + \bigcirc , and

 $(C_eH_s)_sC \rightarrow (C_eH_s)_sC$ + \bigoplus —is it not possible that the electrical conditioning of solution in liquid animonia, and tripliciplinelly l in liquid sulphin disorde, are perfectly analogous phenomena interpretable in similar terms?

An explanation of the conductivity of sodium in liquid in months of the Cody, we and later Franklin and Kiaus, we found that when sodium was dissolved in liquid ammonia, the solution was a good conductor, but there was no visible separation of any poducts of electrolysis at the electrodes, neither were there any signs of polarization. Since conductivity piesupposes the existence of positive and negative ions, perhaps the simplest explanation of the formation of ions

in this case depends upon the change, $Na \rightarrow Na + \bigcirc$, which at once accounts for the positive ions. But what disposition is to made of the negative electron? What is its function? The nitrogen atom in ammonia is tervalent but is regarded as unsaturated since it possesses potentially one free positive and one free negative valence as evidenced by its combination with water

$$(H \ OH)$$
 or hydrogen halide $(H \ X)$ —

$$NH_2 = H_2N_+^{-1} + H_-^{-1}X \Rightarrow HN_-^{-1}$$

Hence, it may be assumed that the negative election (from sodium) neutralizes or polarizes the potential positive valence of the nitrogen atom of ammonia thereby developing a negative valence—

$$Na \rightarrow Na + \Theta$$
,
 $NH_{J}(= H_{J}N\pm) + \Theta \rightarrow NH_{J}$

In other words, the ammonia inolecule is converted into a negative ammonia ion. With positive sodium ions and negative ammonia ions, conductivity is readily explained. Sodium ions

are discharged at the cathode ($\dot{N}a + \bigcirc \rightarrow Na$) and the liberated metallic sodium immediately dissolves and reionizes in the liquid ammonia. The negative ammonia ions are simultaneously discharged at the anode with the immediate regeneration of molecular control of the cont

cular ammonia $(NH_3 + \bigoplus \Rightarrow NH_2)$. This scheme not only accounts for (1) the conductivity, but also explains (2) the absence of products of electrolysis at the electrodes, and (3) non-polarization.

Other facts will now be considered which substantiate the preceding explanation. One of the difficulties encountered in conductivity measurements on solutions of sodium in liquid ammonia is the marked decrease in concentration of sodium through interaction of solute and solvent forming sodamide and hydrogen:—

What is the mechanism of this reaction? The existence of the

ions, Na and NH₃, affords a logical interpretation. Corresponding to the ions in a given solution, there is always the possibility of the existence of some undissociated molecules. In this

instance, union of the ions, Na and NH, would yield the molecule, sodium ammonia, Na.NH, the complete electronic formula being—

It is at once evident that this compound is exceptional in that its nitrogen atom is quadrivalent. Let it now be recalled that just as carbon is tervalent in (C₂H₃)₂C, so nitrogen was shown to be quadrivalent in NO₂, and each of these compounds may be regarded as a free radical. Their reactivity was explained by the abnormal valence of the carbon and nitrogen atoms and their tendencies to revert to the normal valence by gain or loss of electrons. In the same sense, the above sodium-ammona compound may be regarded as a free tadical—an unstable compound containing quadrivalent nitrogen. Greater

^{*}This conception of a free rad-cal embraces the interesting and significant organic amalgams prepared by McCoy and Moore 100 For example, tetramethyl ammonium amalgam was prepared by the electrolysis of cold alcoholic solutions of tetramethyl ammonium chloride using a mercury cathode. In this reaction, the

1 (

stability would ensue if the nitrogen atom became tervalent

through the loss of an election; and, if one of the $\overset{\circ}{H}$ atoms acquires this electron, that hydrogen atom would be rendered neutral, i.e., free or dissociated, and sodium amide would be formed. Combination of liberated hydrogen atoms yields molecular hydrogen. The complete electronic scheme for the formation of sodium and ammonia ions, molecular sodium ammonia, and sodium amide and hydrogen, is as follows:—

$$\begin{array}{c} Na \stackrel{\longrightarrow}{\longrightarrow} \Theta \stackrel{+}{\longrightarrow} Na \\ NH, \stackrel{+}{\longrightarrow} \Theta \stackrel{\longrightarrow}{\longrightarrow} NH_2 \\ \end{array} \rightarrow \begin{array}{c} Na \stackrel{+}{\longrightarrow} -1 \stackrel{+}{\longrightarrow} H \stackrel{\longrightarrow}{\longrightarrow} \\ H \stackrel{+}{\longrightarrow} Na \stackrel{+}{\longrightarrow} -1 \stackrel{-}{\longrightarrow} H \\ \end{array} + H (aH = H_2 = H \stackrel{+}{\longrightarrow} -H),$$

tetramethyl ammonium ion (in which nitrogen is quinquevalent) loyes its positive charge or valence whereupon the nitrogen atom is rendered unad ivalent.

$$(CH_2)_1^+N + \bigcirc \rightarrow (CH_2)_4N$$

(dissolved in Hig) In other words, tetramethyl ammonum, (CH_{ab})N, is a free radical in the same serves that NO, and NaNH, are feer cardicals. But he are to retain the most point of pumbiral and in the properties of the properti

McCoy and Moore state that the tetramethyl ammonium radical acts on solutions of copper and zine salts, replacing these metals in the salts and setting free the metals themselves. The electionic scheme is as follows—

$$2(CH_{2})_{1}N \rightarrow 2(CH_{2})_{1}N + 2 \bigcirc,$$

 $CU + 2 \bigcirc \rightarrow CU.$

These reactions show the striking analogy existing between free radicals and metals, their behaviour being identical in many instances.

Further evidence for the existence of the sodium-ammonia

compound formed by the union of the ions Na and $NH_2 \cap B$ and $NH_2 \cap B$ and $NH_2 \cap B$ solution in liquid ammonia, two molecules of sodium unit with two molecules of ammonia forming $Na_1N_1H_2$. This is readily explained by the existence of the fiee radical, Na, NH_3 in which nitrogen is quadrivalent. The tendency for two such fice radicals to associate would lead to the formation of $Na_2N_2H_3$ according to the following scheme.—

The above reaction is exactly parallel to the union of the two free radicals, nitrogen dioxide, to form nitrogen tetroxide—

or to the union of two free radicals of triphenylmethyl to form hexaphenylethane as previously described.

Having shown that the conductivity of solutions of sodium in liquid ammonia may be explained by the existence of the

ions, Na and NH,, and that their union yields the free radical Na. NH, the instability of this radical in turn explained the formation of NaNH, and of Na₂N₄H₆. The quadrivalent intiogen atom in Na. NH₃ became tervalent in foiming NaNH, and quinquevalent in the compound Na₂N₄H₆. The conductivity of solutions of triphenylmethyl in liquid sulphur dioxide may now be considered.

The triphenylmethyl molecule is converted into a positive on by the loss of an electron (either from the methyl carbon atom or from the para-carbon atom of the quinone ring, quinocarbonium base). This electron converts a sulphur dioxide molecule into a negative ion. This follows from the fact that sulphur dioxide, like ammonia, possesses potentially one free positive and one free negative valence. (Combination with water yields a tautomeric form of sulphurous acid, thus-

$$(80^{\circ} = 0^{\circ}8 \pm) + 11^{\circ}01 \Rightarrow 0^{\circ}8 \xrightarrow{1} H$$

The negative electron from triphenylmethyl polarizes the potential positive valence of the sulphur atom of sulphur dioxide thereby developing a fice negative valence. In other words, the triphenylmethyl molecule becomes a positive ion and the sulphur dioxide molecule becomes a negative ion according to the scheme of the following reactions (1) and (2):—

(I)
$$(C_0H_0)_{\gamma}C \rightarrow \bigcirc + (C_0H_0)_{\gamma}C$$
.
(2) $O_{\gamma}S\pm + \bigcirc \rightarrow \stackrel{-}{SO}_{2}$

The existence of these ions would account for the conductivity of triphenylmethyl in liquid sulphur dioxide solution. The products of electrolysis are triphenylmethyl at the cathode through the reversal of reaction (1), and sulphur dioxide at the anole through the reversal of reaction (2), or the equivalent

change, $SO_2 + \bigoplus \rightarrow SO_2$. This scheme is identical in principle with the scheme for the ionization of sodium and ammonia previously described

Briefly summarized, the interpretations of a number of the characteristic properties of free radicals, particularly triphenylmethyl, in terms of the electronic conception of positive and negative valences, have led to the following conclusions.—

- (1) Free radicals are comparable in many respects to the electrically neutral atomic state,
- (2) The instability, or leactivity, and general chemical properties of free radicals depend upon their development of positive or negative valences through the loss or gain, respectively, of negative electrons. In other words, free radicals may act either as reducing or as oxidizing agents.
- (3) The adaptibility and tendency to lose or to gain elections are properties of certain atoms in the free radicals The valences of these atoms are variable both as to number and to polarity.
 - (4) Not only triphenylmethyl and other triarylmethyls, but

also nitrogen dioxide, sodium-ammonium, tetramethyl ammonium, and other compounds, as well as certain elements, may be regarded as free radicals.

- (5) Some of the peculiar rearrangement reactions manifested by triphenylmethyl are readily interpreted by means of the electronic formula of benzene.
- (6) Evidence is presented in support of the view that there is a close analogy between the electrical conductivity of triphenylmethyl in liquid sulphur dioxide solution, and the conductivity of sodium dissolved in liquid ammonia. In the former instance,

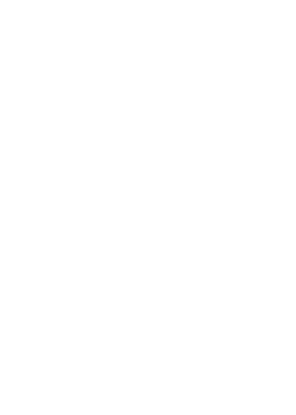
the ions are $(C_0H_3)_{\rm L}^{\circ}$ and SO_2 , in the latter, Na and NH $_2$. The existence of these ions is substantiated by the formation in solution of the free radical, sodium-ammonium, NaVH $_2$, the electronic formula of which lends itself to an interpretation of, and a mechanism for, the reaction in which sodium amide is formed, as well as the compound Na₁N₂H $_2$.

- (7) Finally, it is hoped that the definition and conception of free redicals (herewith developed, applied, and confirmed by interpretations of numerous reactions) may prove to be free from the more or less vague and mystical ideas which have characterized the use of the term "free radical" in the history of chemistry.
- It is only through the employment of the electronic conception of positive and negative valences and electronic formulæ that the interpretations of the existence and properties of free radicals, and the general conclusions here presented, are possible. The literature upon this subject shows that ordinary structural formulæ afford very limited and.



PART III.

PHYSICAL PROPERTIES AND PHYSICO-CHEMICAL PHE-NOMENA MOLECULAR VOLUMES, ABSORPTION OF LIGHT AND FLUORESCENCE



CHAPTED VVII

THE ELECTRONIC FORMULÆ AND MOLECULAR VOLUMES OF

In Part II., Chapters VII. to XVI. inclusive, the constitution of benzene, substitution in the benzene nucleus, and the chemical properties of benzene, as well as many of its derivatives, have been interpreted in terms of the electronic conception of valence and the electronic formula of benzene. The purpose of Part II. has been to substantiate the electronic conception of valence and the electronic formula of benzene chiefly by the enumeration, explanation, and interpretation of chemical properties and physica-chemical Phenomena, such as molecular volume data, light absorption, and fluorescence data to be explained in terms of the electronic conception of positive and negative valence. The discussion and proposed interpretation of these physical and physico-chemical phenomena constitute the subject matter of Part III., Chapters XVIII. for XVIII.

A Molecular Volume Anomalies

Further evidence for the electronic formula of benzene may be found in the development of a relationship between the electronic formulae and the molecular volumes of mono-, di, tri, tetra-, penta-, and hexa-chlorobenzenes. What are the essential features of the relationship existing between the molecular volumes of these six compounds and their respective electronic formulae?

A propos of this question, consider the remarkable anomaly fine noted by Staedel ¹¹ that β -halogen compounds always possess smaller molecular volumes than their isomeric α -halogen compounds. For instance, the molecular volume of CH₃CHXCH₃X is about three units less than that of CH₃CH₂CHX₂. Thus, two structurally similar molecules may possess different molecular

volumes, and there is nothing in the ordinary structural formulæ of these isomers to account for, or to correlate, these differences.

Le Bas 112 has proposed an explanation of some of these anomalies in his hypothesis of partial or incomplete rings, and his examination of the molecular volumes of numerous compounds has led to the following results 112 which must be considered in this paper: (a) If a paraffin compound he substituted by a halogen atom or other unsaturated substituent more than once, the volume of the compound is normal, or subject to the additive rule, provided that the substituents be attached to a single carbon atom. (b) If they are distributed among several carbon atoms, there are always contractions of a constituent nature. It matters not how many hydrogen atoms of the terminal carbon are substituted, the contractions are dependent upon the number substituted in the \$-position (c) For one substitution in the B-nosition the contraction is a little over three units. For two substitutions in the β -position the contraction is about six units, or nearly double the first Substitutions in the y-position always involve greater contractions than substitutions in the B-position . and it is conceivable that substituents in the δ-position would result in still greater contractions.

In explanation of these results Le Bas ¹¹⁴ states that intermediate between open and closed chain compounds are what are
known as partial or incomplete ings "In order that they may
be formed, it is necessary that two atoms or groups, which may
be supposed to possess residual affinity, occur in a hydrocat bon
chain attached to different carbon atoms. If the catbons are
near to each other, there is no need to suppose any variation
from structure usually considered when the tetrahedial arrangement of the valency links of carbon is understood. The additional feature of curvature of the hydrocatbon chain may also be
the normal condution of things In the case of saturated compounds the plane formulæ are figured thus—

If the two attracting groups are united to carbons not in the immediate neighbourhood of each other, we must suppose that they are brought near by the curvature of the hydrocarbon chain, or the alternative supposition just given is true. In any such case, such a structure affects the molecular volumes of comnounds." Thus -

It is not the purpose of this chapter to disparage or disciedit the explanation of Le Bas, but, rather, to propose a new explanation based upon the electronic conception of positive and negative valences as follows. Halogen atoms (and other substituents) may function positively or negatively. Halogen atoms substituted in the a-position alone, or in the 8-position alone. cause no contraction; but each of the halogen atoms in the Bposition occasions a contraction of about three units provided one or more halogen atoms are present in the a-position. May we not assume, then, that the differences or contractions in the molecular volumes of the isomers are due to differences in the polarities of the halogen atoms or substituents, in the a- and B-positions respectively? Further, may we not assume that differences in tolarity are directly related to differences in the relative atomic volumes of the halogen atoms or substituents? For purposes of illustration, suppose that the halogen atoms in the a-position function positively, while those in the β-position function negatively (evidence will be presented shortly) and that the atomic

volume of X is greater than that of X. Then, electronic formulæ,

such as CH,CH,CHXX, and CH,CHXCH,X, correlate the fact that the molecular volume of the a-di-substituted compound is greater than that of the isomeric a. B-di-substituted compound, the difference in the molecular volumes being due to differences

in the atomic volumes of X and X.* Such a correlation cannot be shown by employing the ordinary structural formulæ of the compounds in question.

The above assumptions may now be correlated with a number of facts which are readily interpreted in terms of the electronic

^{*} It should be noted that when different atoms, such as chlorine and carbon, are united, a change in the polarity of one, involving a change in the polarity of the other, would concomitantly alter the atomic volumes of both atoms, but the atomic volume of each atom would not be altered to the same extent. Accordingly, the molecular volume data under consideration in this chapter are regarded as a measure of certain additive and constitutive effects involving the polarity and the atomic volume of certain atoms.

conception of positive and negative valences. It has been noted that when a parafin compound undergoes substitution more than note, the molecular volumes of the substitution products are normal, i.e., subject to the additive rule, provided that the substitutions are linked to a single carbon atom. In this connection consider the direct substitution of chlorine in methane which may be represented thus:—

Successive substitution of positive chlorine atoms would result in the formation of compounds of formulæ.

If we tabulate the molecular volumes (V_m) of these compounds and the volumes of then hydrocarbon groups, the differences between these values will give the volumes of their chlorue atoms which are indicated under (nCI) in the following table, n being the number of chlorine atoms in the respective compounds:—

Compo	und		(V ₂₁₃ .)	Volume of Hydro- carbon Group.	(nCl)		
CH,CI, CHCI, CCI,	:	:	65 I 84 5 103 7	14 8 18.2 14 8	1 × 55 7 3 × 55 0 5 × 51.2		

The data show a slight increase in the atomic volumes of the chlorine atoms as they accumulate in the molecule. The electronic schemes for the substitution reactions (in the absence of intramolecular oxidation-reduction reactions) indicate that the substituted chlorine atoms function positively. The molecular volume data indicate that the average atomic volume of these positive chlorine atoms is very nearly equal to 220.

The next question to be considered naturally relates to the adomic volume of the chlorine atoms which function negatively. An extension of the hypothesis to the constitution and molecular volumes of chiylene and ethylidene dichlorides leads to some interesting conclusions. The values with respect to these isomers are as follows—

Compound	(V _m)	Volume of Hydro carbon Group	(#C1)		
CH, CHCl, .	88 g	44 5	2 × 22.2		
CII,CI CH2CI .	85.3	41'5	2 × 20 4		

From the data it appears that the average atomic volume of the chlorine atom in ethylidene dichloride is 22.2: in ethylene dichloride, 20.4. Correctly to interpret this constitutive effect, account must be taken of the fact that in the majority of the aliphatic compounds of the type RCl in which only one chlorine atom is substituted on a carbon atom, the average atomic volume of this single chlorine atom is 21'5. Since the chlorine atoms in ethylene dichloride are singly attached to separate carbon atoms and their total volume is 40.8, the volume of each cannot be 21.5 Consequently, it follows that if one of these chlorine atoms has the usual volume of 21 5, then the volume of the other chlorine atom must be 103, ie, the difference between their total volumes, 40.8 and 21.5. While none of these values is absolute, the data indicate a possible correlation of molecular volume data and electronic formulæ. Thus, if the α-chlorine atom has a greater atomic volume than the B-chlorine atom and functions positively, then, in contra-distinction, the \(\beta\)-chlorine atom would be assumed to function negatively. The correlation of the electronic formulæ and atomic volumes of the chlorine atoms of ethylidene and ethylene dichlorides may be summarized thus :-

From this point of view, the difference between the molecular volumes of ethylidene and ethylene di-chlorides is not dependent on the idea of the formation of a partial ring, as assumed by Le Bas in the case of ethylene dichloride, but is due to a difference in the atomic volumes of the a- and β -chlorine atoms which are assumed to function positively and negatively, respectively Further evidence for the assumption of this difference of polarity is found in the union of chlorine with ethylene. The hydrolysis of chlorine (previously indicated) vielding hydrogen chloride (H. Cl) and hypochlorous acid (HO . Cl) indicates that the follorine molecule may yield Cl and Cl on dissociation Accordingly, the addition of chlorous to ethylene is represented thus—

The "opening" of the double bond of ethylene may be regarded as the simultaneous development of a free positive valence on the one carbon atom and a free negative valence on the other, which valences, respectively, bind the negative and positive chlorine atoms. In other words, the chlorine atoms in the a- and B-positions are of opposite polarity.

An analogous situation, indicating that the a-substituent tends to function positively while the β -substituent is negative,

is found in the addition of hydrogen chloride (\hat{H} . \hat{Cl}) to acrylic acid yielding β -chloropropionic acid exclusively, $^{\prime}$ thus —

Note also in this connection the addition of hydrogen bromide to a-bromo acrylic ester. Again the positive substituent unites with the a-carbon atom while the negative substituent assumes the B-position, thus —

This compound, a, β -di-bromopiopionic ester, is also the product of the addition of bromine (Br. Br) to acrylic ester, and in

and contrafter (C===C) (Fry, Z fhysil. Chem., 76, 400 (1911)), renders possible an interpretation of many of the anomalous addition reactions studied by Michael from the standpoint of his "chemical neutralization" theory

^{*} Addition reactions involving the two types of double bonds, diflex (C $\stackrel{+}{=}$ -C),

view of the preceding facts the a- and β -bromine atoms are, respectively, positive and negative. Now the molecular volume of a, β -th-homopropionic ester is 39 units less than that of its isomei, the a, a-compound. Thus it appears that the a-substituents have a greater atomic volume than the β -substituents and a relationship exists between the electronic formulæ of these isomeies and their molecular volumes which is summarized as follows —

The preceding facts and their correlation with the electronic formulae lead to the assumption that the atomic volume of the positive halogen substituent is greater than that of the negative halogen substituent. Consequently, the differences between the molecular volumes of the isomers in question is related in some way (to be considered later) to differences in the polarities of the substituents in the a- and \(\theta\)-positions. This hypothesis may now be extended to the explanation of the remarkable anomalies met with in the atomic volumes of the chlorine atoms in the several chlorobenzenes, especially hexa-chlorobenzene. The existence of a relationship between the molecular volumes of the chlorobenzenes and the electronic formula of benzene will be definitely indicated.

B. The Electronic Formulæ of the Chlorobenzenes.

In previous chapters numerous facts have been presented and interpreted by means of the electronic formula of benzene. A substitution tule has been developed showing that when substituents are of the same sign or polarity they will occupy positions which are meta to each other, but if two substituents are of opposite sign or polarity they will occupy either ortho or para positions to each other. The development of this rule depended upon the fact that the electronic formula of benzene is the only formula which presents conjointly a structural basis and an electronic interpretation of the relations between ortho and para positions and substituents in contradistinction to meta positions and substituents. Now, if the electronic formula of benzene

is subject to correlation with the molecular volumes of the chlorobanzenes, the following conditions must be fulfilled: (1) Substituted chlorine atoms occupying ortho or para positions to each other must be of opposite polarity while those occupying meta positions to each other must be of the same polarity (2) It has been indicated that chlorine atoms which function positively possess a greater atomic volume than those which function negatively. Consequently, positive chlorine atoms in the benzene nucleus must possess greater atomic volumes than the negative chlorine atoms in the corresponding para or ortho positions. These conditions would be fulfilled completely if it is shown that, for instance, in hexa-chlorobenzene

the positive chlorine atoms in positions 1, 3, and 5 possess greater atomic volumes than the negative chlorine atoms in positions 2, 4, and 6 To this end, consider, first, the electronic schemes for the formation of the six chlorobenizenes whose molecular volumes present the remarkable anomalies that demand an explanation.

The direct substitution of chlorine in the benzene nucleus, generally effected with the aid of halogen carriers, corresponds to the following simplified scheme I.—

in which positive chlorine is introduced. The extended substitution of chlorine yields the several di-, tri-, teta-, penta-, and hox-a-chlorobenzenes. Direct substitution, as indicated, effects the introduction of a positive substituent, but the electronic formula of benzene requires that poly-substituted derivatives embody negative substituents as well. The existence of the negative substituents depends upon the electronic tautomersing (for ex-

ample, $C_6H_5Cl \Rightarrow C_6H_5Cl$) of some of the derivatives which in turn undergo further substitution With reference to the formation of para-di-chlorobenzene, or any other di-substituted derivative. recall that if a given substituent in the benzene nucleus is positive, the entering positive substituent will occupy the meta position, but if the mono-substituent is negative, the entering positive substituent will occupy para or ortho positions. The chloringtion of mono-chlorobenzene yields chiefly para-di-chlorobenzene, Consequently, electromer C.H.Cl (and not electromer C.H.Cl) undergoes substitution according to scheme II. .-

Scheme II.

$$\vec{C}$$
 \vec{H}
 \vec{H}

Schemes I, and II, in conjunction with the principle of electronic tautomerism show that a given positive substituent may, under certain conditions, function as a negative substituent, but in para-di-chlorobenzene the halogen atoms are of opposite polarity. This not only follows from the electronic formulæ but is confirmed by experimental facts, notably the reactivity of the negative chloring atom and the non-reactivity of the positive chlorine atom when para-di-chlorobenzene is heated with methyl alcohol solution of sodium methylate. The reaction conforms quantitatively to the electronic equation .-

$$\overset{+}{\text{Cl.}} \cdot \text{C}_0 \text{H}_4 \cdot \overset{-}{\text{Cl.}} + \overset{+}{\text{Na.}} \cdot \text{OCH}_3 \ \rightarrow \ \overset{+}{\text{Cl.}} \cdot \text{C}_0 \text{H}_4 \cdot \text{OCH}_3 \ + \overset{+}{\text{Na.}} \cdot \overset{-}{\text{Cl.}}$$

Another remarkable difference between the chloring atoms in para-di-chlorobenzene will shortly be considered in connection with their atomic volumes.

In the complete scheme for the formation of the six chlorobenzenes in question, a number of systems of electronic tautomerism are involved, but in the following abbreviated scheme III. only those electronic formulæ have been included which are directly related to the present investigation, namely, the correlation of the molecular volumes of mono-, 1:2 di-, 1:2 4 or 12 *

1 4:5 tti, 1 2:4 5 tetra, penta, and hexa-chlorobenzenes with their respective electronic formula. The electronic formula of these compounds are indicated in the following scheme III by the letters A. B. C. D. E. and F:—

$$\begin{array}{c} \prod_{i=1}^{H} \prod_{j=1}^{H} C_{i,j} \\ \prod_{i=1}^{H} \prod_{j=1}^{H} C_{i,j} \\ \prod_{i=1}^{H} \prod_{j=1}^{H} \prod_{i=1}^{H} \prod_{j=1}^{H} \prod_{j=1}^{H} \prod_{i=1}^{H} \prod_{j=1}^{H} \prod_{$$

The scheme involves three systems of electronic tautomerism, namely, $(A \rightleftharpoons A')$, $(C \rightleftharpoons C')$, and $(E \rightleftharpoons E')$. Beginning with benzene, each of the six successive chlorinations introduces only

one positive substituent, Cl, as has been illustrated in preceding schemes I, and II Note that A', C', and E' (the respective electromers of A, C, and E) yield on chlorination B, D, and F, respectively. Compounds B, D, and F do not possess electromers because they contain an even number of symmetrically substituted chlorine atoms. On the other hand, the electromic formulae A, C, and E are in tautomicric equilibrium with their respective electromers A', C', and E' since they embody an uncirce number of substituted chlorine atoms and the number of their positive substituents is unequal to the number of their negative substituents.

Before tabulating the molecular volume data, it will facilitate comparisons if the electronic formulae of the six chlorobenizenes are rewritten in juxtaposition and with the positive substituents in positions 1, 3, and 5, and the negative substituents in positions 2, 4, and 6. Such an arrangement will not alter in the least the electronic formulæ of these compounds as derived in the preceding scheme the relative positions and polarities of the substituents remain the same

C. The Molecular Volumes of the Chlorobenzenes.

The preceding arrangement of the electronic formulæ of the six chlorobenzenes (A-F) serves as a key to the following table of molecular volume values :-

Chlorobenzenes		V _{ex}	V'm	V _e CI	Δ
A Mono B I; 4-Di-	(odd) (even)	114 6 130 9	g2 8 8q 6	21 S 41 3	19'5
C 1.4:5-Tri-	(odd)	149.1	86 4	62 7	21'4 18 9
D. 1 2.4.5-Tetra-	(even)	164.8	83 2	81.6	223
E 1 2 3:4.5-Penta- F, 1.2 3:4 5 6-Heva-	(odd) (even)	183.9	80 0 76 8	103,0	19:3
1.1.2 3 4 5 Olieta	(even)	2000	700	123.2	

The first column embodies the names of the six chlorobenzenes, A to F, inclusive The designation (odd) or (even) refers to the number of chlorine atoms in the respective compounds. The former are unsymmetrical in structure, the latter, symmetrical. The molecular volumes of the chlorobenzenes, originally determined by Jungfleisch,115 and subsequently retabulated by Le Bas, are given in the second column, V., third column indicates the molecular volumes of the hydrocarbon groups of the compounds, V'... The difference between the molecular volume of a given chlorobenzene and its hydrocarbon group (Vm - V'm) gives the molecular volume of its chlorine

atoms, V.Cl. column four. The last column is one of differences (A) between the total volumes of the chloruse atoms of the successive compounds For example, the atomic volume of the chloring atom in mono-chlorobenzene is 21%. The combined atomic volumes of the two chlorine atoms in 1 · 4 di-chlorobenzene is 41'3. The difference between these values, namely, 19:5, is, therefore, the volume of one chlorine atom while that of the other in the para position is 21 8. The electronic formula of benzene has shown that substituents para to each other are of opposite polarity. Since previous investigations have indicated that the positive chlorine atom may have a greater atomic volume than the negative chlorine atom, the correlation between the atomic volumes and the electronic formula is maintained in paradi-chlorobenzene. Further correlations are maintained throughout the series of the six chlorobenzenes since the column of differences (A) shows that the chlorine atoms successively substituted vary in their atomic volume values by an average difference of ± 2 0 units This simultaneous variation in polarity and atomic volume is emphasized by the retabulations in the following table :--

	Relative Position	Polarity	Atomic Volume (Positive)	Atomic Volume (Negative)
A 1st chlorace atom . B 2nd	(1) (4) (5) (2) (3) (6) Mean	(+) (-) (+) (-) (+) (-) values	21 8 21 1 22 3 21 1 21 1	19'5 18'9 19 3

It is, therefore, evident that a marked difference (average $\Delta=2.6$) exists between the atomic volumes of the chlorine atoms in positions 1, 3, and 5 and those in positions 2, 4, and 6; and, since the electronic formula of benzene and the substitution in the as shown that substituents in positions 1, 3, and 5 are opposite in polarity to those in positions 2, 4, and 6, it follows that a definite relationship exists between the electronic formulae of the xix oblivation exists and their respective molecular

additional evidence, from a physical point of view, for the electronic formula of bensene.

Various facts presented in this chapter seem to indicate that a change in the polarity of a halogen atom causes a variation in its atomic volume. Any attempts to explain this in the present state of our knowledge would plunge us fatally into the metaphysics of an atomic structure maelstrom. Notwithstanding the incomplete knowledge of the constitution of the atom as manifested by the many different hypotheses relative to the nature and disposition of "valence elections," it is quite conceivable that variations in the relative positions of the valence electrons which determine the polarity of an atom may likewise cause variations in the atomic volume of the atom. This assumption may lead to an explanation of other anomalies in molecular volume relationships. For instance, in the compounds methylene dichloude, chloroform, and carbon tetrachlouide, previously noted. the average atomic volumes of the chlorine atoms are 21.5. 22 0, and 22'2, respectively. This small increase in the average volumes of the chlorine atoms as they accumulate about a given carbon atom may be due to the relative positions of the valence elections between the carbon and chlorine atoms which may mutually alter the respective atomic volumes of these atoms. It has also been noted that for one substitution in the 8-position of a compound the contraction in molecular volume is a little over three units while substitutions on the v- and δ-positions result in even greater contractions. These anomalies may also be related to differences in the relative positions of the valence electrons between the carbon atoms and the halogen atoms substituted in the a-, β -, γ -, and δ -positions, thus causing varying differences in the degrees of contractions.

In conclusion, a quotation from Le Bas should be noted --

"There is no doubt that in spite of the care taken, many parts of the present theory of molecular volumes may have to be altered later as data accumulate, and as our knowledge of the physical property increases The identification and explanation of constitutive effects is not always easy. Some particular atomic values—generally those found in the homologous series R.—X are taken as standard, and by the method of summation the value $\Sigma_n V_{nl}$ found. The difference, $V_m \Sigma_n V_m$ then measures the constitutive effect. Sometimes a mean atomic value is

taken, and it then follows that no account is taken of variations. The great difficulty is to identify the effect with a particular atom or group. When this seems possible, it sometimes happens that other atoms or groups might equally well be identified with the effect in question. Only a careful examination of a lattre number of data can overcome these difficulties. It will generally be found that the constitutive effects are traceable to some modifications in particular atomic values, and a considerable advance is made when we are able to ascertain for certain which atoms are marked by the variation in question and by how much "

Apropos of the above quotation, and as a summary of the present chapter, it has been shown that -

- (1) Halogen atoms which function positively appear to possess different atomic volumes from those which function negatively Consequently, it is possible to correlate certain additive and constitutive effects apparent in the molecular volumes of certain compounds with their electronic formula. These effects cannot be explained by means of the ordinarily employed structural formula.
- (2) Definite relationships exist between the molecular volumes of six different chlorobenzenes and their respective electronic formula. These relationships further confirm the electronic formula of benzene
- (3) It is suggested that any variations in the relative positions of the valence electrons which determine the polarity of an atom may likewise cause variations in the atomic volume of the atom

CHAPTER XVIII

DYNAMIC FORMULÆ AND THE ULTRAVIOLET ABSORPTION SPECTRUM OF BENZENE.

A. Colour in Relation to Chemical Constitution.

It is the purpose of this chapter to present the development of a relationship, in the nature of a linear function, between the systems of dynamic equilibria of the various electromers of benzene and the oscillation frequencies of the seven bands which characterize the ultraviolet absorption spectrum of benzene solutions. The existence of such a relationship would not only further substantiate the electronic formulæ of benzene but also afford a new explanation of colour in relation to chemical constitution.

Watson, ¹⁶ in the pieface to his recent monograph, "Colour in Relation to Chemical Constitution," states that "the early theories as to the relation between colour and constitution, such as the quinomoid theory and Nietzki's rule, have proved of great value for practical purposes, viz., in the production of dyestuffs and especially for the preparation of dyestuffs of any required shade; but more scent researches have shown that these classical theories are by no means adequate. Modified and new theories have been proposed which agree better with the known facts. The quest of the altimate anxe of colour has revealed the great complexity of the problem and has shown the need for further work in this direction."

The fundamental cause of the absorption of light by benzene, as proposed in this chapter, is necessarily complex since it has to deal with the somewhat complicated systems of dynamic equilibria of the electromers of benzene. But if the absorption of light of known oscillation frequency can be correlated with the electronic formulae of benzene, then it may be concluded

that some definite advance has been made in the quest of the ultimate cause of colour in its relation to chemical constitution.

B. The Systems of Dynamic Equilibria of the Electromers of Benzene.

A statement made by Stewart ¹¹⁷ should be recalled at this point "It is becoming generally recognized that the benxeene molecule is in a state of continual vibration, and that the only satisfactory space formula will be one which represents all the other formulae as phases of its own motions, and which may even suggest the possibility of new phases as yet unrecognized. The main outlines of such a formula have been indicated by Collie, and it seems probable that any space formulae of benzene which may be proposed in the future will agree with his in essentials."

The plane projections of Collie's space formulæ and the twofold movements which are attributed to the system have been described and illustrated in Chapter VIII, Section B (p. 48). Collie's space formulæ are correlated readily with the Keknië and centric formulæ since, through their movements, they are mutually interconvertible.

Again, recall that a complete application of the electronic conception of positive and negative valences to the benzene molecule, involves five types of carbon atoms, namely,

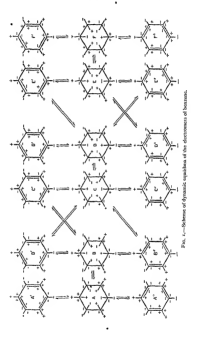
and if beitzene nuclei (centife formulæ) are composed of these several types, each nucleus consisting of three pairs of the combined types I and V., II, and IV., and III. and III. symmetrically co-ordinated, six and only six centric electronic formulæ (electromers) are possible, as previously shown in Chapter VIII

Now, if in place of the one centric phase of Collie's space formula, there be sax centric electromers, then there will be six times as many Kekulé and other phases in a complete system of dynamic equilibria of the electromers of benzene. Before this extended system is developed, consider specifically the plane projections of those electromers that are related to the centric formula which is composed of carbon atoms of type III. —

Note that the first and second Kekulé phases present two distinct types of double bonds In the former, each of the two bonds (Faraday tubes) comprising a double bond has the same direction, since their adjacent ends are of the same sign or polarity, In the latter, the bonds have opposite direction, since their adjacent ends are of opposite sign or polarity. Hence these two types of double bonds may be designated by the terms diplex and contraplex respectively.

In the complete scheme of dynamic equilibria of the electromers of benzene, presented in Fig. 1, p. 188, the first and last phases, which are respectively related to the first and second Kekulé phases, have been omitted for the sake of brevity. The centric electromer A, the starting-point of the scheme, so to speak, is composed of carbon atoms of the types I. and V. Electromer A functions as the intermediate phase between A' and A".

Kekulé held that the atoms of carbon oscillate in the molecule of benzene in such a way that A' would represent the constitution of the molecule at one period of oscillation, and A" the constitution at another period. This conception introduces a kind



of structural modification differing both from ordinary isomerism and tautomerism for which Bruhl 118 has suggested the term phasotropism, in the sense that "the unaltered benzene nucleus and analogous ring systems are phasotropic". On the electronic basis, A' and A" may be termed phasotropia electromers, neither of which could revert to the centric electromer A, unless there was a rearrangement of bonds or Faraday tubes. Such a change would result in another state of equilibrium in the centric electromer or phase.

Faraday introduced the term "dielectric polarization" to describe the condition of a non-conductor or dielectric, as he conceived it, when in a state of strain under the action of two adjacent charges of positive and negative electricity, as, for example, in the condenser. Accordingly, the centric phase may be assumed to present a state of strain, or of meta-stable equilibiium, by virtue of the balance between three positive and three negative charges within the ring; hence it may be said to be in a condition of dielectric polarization. The rearrangement of the three positive and three negative charges of the centric electromer in such a manner that the positive charges occupy the positions antecedently held by the negative charges, and vice versa, produces at one and the same time not only a similar condition of dielectric polarization, but also another centure electromer Thus, the centuc electromer A composed of carbon atoms of the types I and V, undergoes centric rearrangement yielding another centric electromer B composed of carbon atoms of the types II. and IV. There are three distinct centric rearrangement equilibria, or transitions, in the complete scheme; namely, A = B, C ≠ D, E ≠ F. Note that centric rearrangement is structurally impossible between B and C, and between D and E.

In Collie's space formulæ for benzene, the centric phase is intermediate between two, and only two, Kekulé or phasotropic phases. In the proposed scheme, in which every electronic formula is the plane projection of a space formula, a given centisc electromer may be the intermediate phase between two, three, or four phasotropic electromers. For example, A is the intermediate phase between A' and A"; B, between B', B", and C'; C. between B', B", C', and C" Note that the transition from B to C may be effected through B', B", or C', but not through C; also, the transition from D to E may be effected through E', E".

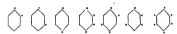
and D", but not through D'. In this manner the inter-relations of the eighteen electromers comprising the complete scheme of equilibria may be readily traced.

The scheme as thus outlined, presents, when considered dynamically, two types of dynamic equilibria which shall be designated as primary and scoulary yxtems of phasotropic equilibria. A primary system or transition involves three phases: namely, one centre electromer and any two of its directly related phasotropic electromers, as for example, the transitions $A' \approx A \approx A'$; $B' \approx B \approx B''$; or $B' \approx C \approx C''$. A secondary system or transition involves four phases; namely, two centric electromers, which are interconvertible by centre rearrangement, and one phasotropic electromer of each of the two interconvertible centric electromers, as for example the transitions of $A' \approx A \approx B \approx B''$, or $B' \approx C \approx D \approx E''$. Twenty primary transitions and twenty-eight secondary transitions are embodied in the complete scheme. They will be tabulated and discussed later.

The possibility of correlating these various electromers and their systems of dynamic equilibria with the oscillation frequencies of the seven bands of the absorption spectrum of benzene constitutes the present problem. Balv and Desch 110 maintain that the benzene absorption bands are to be accounted for by the synchronous oscillations of the benzene molecule in some such way as a tuning-fork vibrates in response to a note of definite pitch. In terms of the electron theory, they state that "combination between two atoms is accompanied by the passage of one or more electrons from one atom to the other producing one or more Faraday tubes of force between them, each Faraday tube representing the chemists single bond. If by some means we cause the rearrangement of these linkings or Faraday tubes, it is clear that there must occur a vibiational disturbance in the systems of electrons of the atoms concerned. Now we have direct evidence of these disturbances in the fluorescence of tautomeric substances as shown by Hewitt 126 It is, therefore, only natural that the converse takes place, namely, the absorption of light by tautomeric substances."

Baly and Desch show that this process in the aliphatic compounds involves the make and break between a carbon and an oxygen atom, thus—

the stars being attached to those atoms which are undergoing the linking change Compounds which manifest keto-enol tautomeusm display an absorption band. Applying this idea to benzene, Baly and Collie 121 differentiate between the transition phases in which any pair of carbon atoms, or any four, or all six are concerned, the various transition phases being represented as follows -



These seven forms are held to represent all the possible conditions of making and breaking the linkings which can occur. each per se being the origin of a separate absorption band

This hypothesis of Balv and Collie does not offer any definite explanation as to how the seven phases of benzene may be brought about, apart from being the result of some form of motion of the atoms in the molecules. Moreover, since it does not attribute or relate any one of the above seven phases to any one of the seven absorption bands, it thereby fails to indicate any possible numerical relationship between the various phases and the oscillation frequencies of the seven bands. An hypothesis designed to meet these deficiencies is offered in the following paragraphs.

C. The Electronic Interpretation of Keto-Enol Tautomerism in relation to the Absorption of Light.

Accepting the finding of Baly and Desch that an absorption band is the result of coexistent ketonic and enolic forms in dynamic equilibria, three questions arise ---

- (1) What is the significance of the keto-enol condition of dynamic equilibrium from the standpoint of the electronic formulæ of keto-enol compounds?
 - (2) Is this significance common both to the keto-enol systems

of dynamic equilibria and to the primary and secondary systems of phasotropic equilibria of the electromers of benzene?

- •(3) How are the systems of phasotropic equilibra to be correlated. (a) with the existence of seven bands in the absorption spectrum of benzene, and (b) with the oscillation frequencies of each of the seven bands? The following answers are proposed ---
- (1) The electronic formulæ of ketonic and enolic modifications are derivable from the electronic formula of one of the five typical compounds (see Chapter III, Section B, p. 17), namely, formaldehyde. The formula of an aldehyde or of a ketone is indicated by replacing one or two hydrogen atoms, respectively, of formaldehyde by alkyl radicals -

Keto-enol tautomensm involves the migration to and fig of a hydrogen atom between carbon and oxygen atoms. Having indicated the electionic formula of a ketone, the keto-enol tautomeric equilibrium may be represented electronically as follows:-

That portion of the above scheme within the brackets illustrates the "opening up" of the dipler double bond between the carbon and oxygen atoms, and the ionization of the migrating hydrogen

ion, H. This intermediate condition presumably precedes the formation of the enolic modification which embodies a contraplex double bond between the two carbon atoms,

Note particularly that the rearrangement of valences or Faraday tubes involves a change in the nature or type of double bonds as found in the keto- and enol-forms. The former presents a diplar double bond between the carbon and the oxygen atom; the latter, a contrapler double bond between the two carrion atoms. Hence the electronic significance of the keto-enol transition resolves itself to a condition of dynamic equilibrium involving transition from diplex to contraplex double bonds and vice versa. Systems of dynamic equilibria involving changes from diplex to contraplex double bonds and vice versa. When the contraplex double bonds and vice versa, will be termed contraplex-diplex transitions. This disposes of the first question.

(a) With reference to the second question, the hypothesis now proposed does not limit contraplex-diplex transitions to keto-enol tautomerism, but further assumes that the occurrence of contraplex diplex transitions in any types of dynamic equilibria constitutes the structural and the electronic crylanation of absorption bands, that is, of colour. The occurrence of these transitions among the primary and secondary systems of phasotropic equilibria of the electromers of benzene have been noted; hence, they naturally constitute the basis of the explanation of the absorption spectrum of benzene. The discussion of the third question now follows

Since there are seven absorption bands in the benzene spectrum, and it is assumed that the transition from contraplex to diplex bonds and vice versā, is the electronic explanation of an absorption band, then there must be seven such distinct systems involved in the complete scheme of phasotropic equilibria. A tabular arrangement (see next page), (cf. Fig. 1, p 188) of all the possible primary and secondary systems of phasotropic equilibria presents twenty of the former and twenty-eight of the latter.

The electromers containing diplex bonds are \tilde{A}' , A'', C'', D', F', and F', while those containing contaplex bonds are B', B', C', D', E', and B''. Only those equilibria, either primary or secondary, which involve transitions from an electromer containing diplex to one containing contaplex bonds, or vice versa, are followed by asterisks These only may function as the origin of the absorption bands.

Since six centric electromers figure in the complete scheme of equilibria, there are, accordingly, six classes of primary transitions, each class involving one centric phase. Only six of the twenty primary transitions are asterisked, namely, those ^

	,
Primary Systems of Phasotropic Equilibria,	Secondary Systems of Phasotropic Lquibbria
{ 1. A' - AA" ∫ 2. B'-B-B"	t. A'—A—B—B' * . A'—A—B—B' * 3 A''—A—B—B' * 4 A''—A—B—B' *
3 B'-B-C' 4 B"-B-C'	5. A'—A—B—C' * 6. A"—A—B—C' *
5. B' C B" 6 B' C C' B" C C'	7 B'-C-D-D' * 8. B"-C-D-D' * 9 B'-C-D-D' 10. B"-C-D-D"
7. B" - C - C" 8. B" - C - C" 9. B" - C - C" 10. C' - C - C" * A	11. B'—C—D—E' 12 B'—C—D—E' 13 B"—C—D—E' 11. B"—C—D—E'
\[\begin{pmatrix} \text{ii} & \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	15. C' \$C-D-D' * 16. C' -C-D-D' 17. C''-C-D-D' 18. C''-C-D-D' 18. C''-C-D-D'' 19. C''-C-D-D''' 19. C''-C-D-D'' 19. C''-D''-D''-D'' 19. C''-D''-D''-D''-D''-D''-D''-D''-D''-D''-
14. D"—D—E" 15. D"—D—E" 16. F'—D—E"	19. C' —C—D—E' 20. C' —C—D—E' 21. C''—C—D—E' 22. C''—C—D—E'
17 E'-E-F" 18 E'-E-D"	23 D"-E-F-F" * VI.
(20 F'-F-F"	25 E'-E-F-F' 26. E'-E-F-F' 27. E''-E-F-F' 28 E''-E-F-F''

numbered 8 to 13 inclusive, which in turn are subdivided into four distinct groups as follows -

```
Group (a) 2 transitions, numbered 8 and 0.
  ,, (β) I transition,
,, (γ) I ,,
(δ) 2 transitions,
                                 .,
                                        To.
                                         12 and 13
```

Since the complete scheme involves three centric rearrangements, A & B, C & D, and E & F, there are accordingly three classes of secondary transitions. They are numbered 1 to 6, 7 to 22, and 23 to 28 inclusive, respectively Eighteen of the twenty-eight secondary transitions are asterisked and fall naturally into servi distinct groups as follows .-

Group		4	transitions,	numbered	1	to	4 inclusiv
	П	2	,,	,,	5	ລກ	d 6,
**	HI.	3	12		7	,,	8,
,.	IV	2	11	**	15	•1	18,
**	ν,	2	13	₹"	21	• • • • • • • • • • • • • • • • • • • •	22,
2.0	VI.	2	11		23	"	24,
**	VII.	4	1)	22	25	to	28 inclusive

An eighth group, transitions II to I4 inclusive, does not embrace any contraplex-diplex equilibria, consequently, it plays no part in the development of the present hypothesis.

D. The Correlation of Oscillation Frequencies and Contraplex-Diplex Transitions.

The existence of seven distinct groups of scoundary contraplicadiplica systems of equilibria constitutes the basis of the correlation of the seven absorption hands of benzene. The oscillation frequencies of the heads of these bands as determined by Baly and Collie are as follows:

Band			Oscillata
One			. 37-25
Two			3765
Thice			3830
Four			3915
Five			4025
Six .			1110
Seven			4200

In attempting to correlate the seven groups of secondary contraplex-diplex equilibria with the above oscillation frequencies. let it be assumed that the number of transitions in each of the seven groups functions successively and collectively in the production of the seven bands. Naturally two, the smallest number of secondary transitions contained in a single group, may then be assumed to function in the production of the band of lowest frequency; namely, band One, 3725. On the other hand, since there are altogether twenty-four contraplex-diplex transitions in the complete system, it may likewise be assumed that all of these function in the production of band Seven of highest frequency. 4200. In other words, the vibrations of two secondary contraplex-diplex transitions are synchronous with light waves of frequency 3725, while the vibiations of twenty-four (i.e., eighteen secondary plus six primary) contraplex-diplex transitions are synchronous with light waves of frequency 4200

These assumptions relative to the origin of bands One and Seven are of no merit or value unless some means is at hand of determining the numbers of transitions involved in the production of each of the five remaining bands. Furthermore, these numbers must be whole numbers since the present hypothesis requires that they be the sums of the transitions in the secondary groups. I. to VII. inclusive, and the transitions in the primary groups,

 α , β , γ , and δ . In other words, a linear function must be shown to exist between the oscillation frequency of each of the seven bands and an integral number of contraplex-diplex transitions. The possibility that the frequencies of the absorption bands of benzene are a function of a series of whole numbers is suggested by the discovery of Balmer, ¹²⁴ that the wave-lengths of the lines in the hydrogen spectrum are a function of successive whole numbers. His formula:

$$\lambda = h \frac{m^2}{m^2 - 4} \cdot 10^{-6}$$

in which h = 3646·13, and m is given the values 3, 4, 5, 6, etc., is only one of several showing the existence of similar series in the line spectra of various elements

A similar relationship becomes apparent if the oscillation frequency 3725 (band One) involving two transitions and 4200 (band Seven) involving twenty-four transitions serve as two points in a system of rectangular co-ordinates, namely (3725, 2) and (4200, 24) The calculated equation for a straight line passing through these two points is v = 21.591x + 3681.818in which (r) is the oscillation frequency of a given band and (x) is the number of contraplex-diplex transitions functioning as the origin of the given band. Assuming on the one hand that the frequencies (y) as determined by experimental observation are absolutely correct, the corresponding numbers of transitions (x) involved therein may be calculated. On the other hand, employing as values for (a) those whole numbers which are most closely approximated by the previously calculated values of (x), the corresponding values of (y) may in turn be ascertained. Thus theoretical and actual values may be compared. In Table No. 1-TABLE No r

(1 = 21 59T1 + 3681 818)

A	В	(A - B)	D,	E	(D - E)
2 000 5 553 6 863 to 860 15 895 19 832 24 000	2 1 7 11 16 20 24	0°000 - 0°147 - 0 137 - 0 200 - 0 105 - 0 168 0 000	3725 3765 3830 3915 1025 4110 4200	3725'000 3768'181 3832'951 3919'318 1027'2'2 4113'536 4200 000	0 000 - 3'181 - 2 954 - 4'318 - 2'272 - 3 636 0'000

column A contains the calculated values for (r): column B, the series of whole numbers most nearly approximated by the calculated values of column A; column C, the differences between values in columns A and B: column D, the oscillation frequencies as experimentally determined by Baly and Collie; column E. the calculated frequencies (y) corresponding to the assumed whole number values for (a) in column B, column F, the differences between the actual frequencies in column D and the calculated frequencies of column E

The values in column A approximate very closely the whole numbers in B as is evidenced by the column of differences, C. This favours the assumption that the oscillation frequencies are a function of whole numbers. Furthermore, the frequencies in column D calculated upon this assumption approximate the actual frequencies in column E as is evidenced by the column of differences F, which are practically within the limits of error of experimental observation.

The calculated values of Table No. 1 are based upon the assumption that the frequencies of bands One and Seven are absolutely correct while the frequencies of the intermediate bands were not considered in determining the equation of the line relating frequencies and the numbers of contraplex-diplex transitions. Numerical values of greater significance are to be found in another equation, namely, v = 21.6063x + 3679.296, which is derived by an application of the method of least squares to the whole numbers (x) and the actual oscillation frequencies (v) taken from columns A and D of Table No. 2.

TABLE No. 2. (y = 2160631 + 3679296)

Α.	B (B -	A) D	E	(D - E.)
7 11 16 20	2 115	034 3765 025 3830 091 3915 000 4025 066 4110	3722'509 3765'721 3830'540 3916 965 4021'996 4111 121 1197 847	+ 2 (91 - 0 721 - 0'540 - 1 965 + 0 604 - 1 421 + 2'153

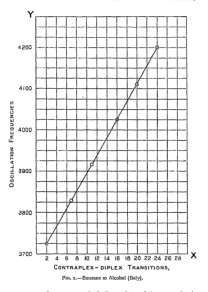
The deviations, column C, of the calculated numbers, column B, from the assumed whole numbers, column A, are so slight as fully to warrant the assumption that the oscillation frequencies of the seven bands, column D, are a linear function of the corespending whole numbers, column A, which are the numbers of the contraplex-diplex transitions involved therein. This hypothesis is further substantiated in that the deviations, column P, from the experimentally determined values, column D, are well within the limits of error of experimental observation. The probable error in the value of any of the oscillation frequencies, as calculated by Peters' formula, is only 17328.

This remarkable relationship between the series of seven whole numbers and the oscillation frequencies of the seven absorption bands of benzene (columns A and D respectively of Table No. 2) is graphically illustrated in Fig. 2. The seven whole numbers (X-axis) and the corresponding oscillation frequencies (Y-axis) permit the location of seven points in the system of rectangular co-ordinates. A line drawn through each of these seven points is practically a straight line. This, in conjunction with the data of preceding Tables I and 2, both warrants and substantiates the hypothesis that the oscillation frequencies of the heads of the absorption bands of benzene are a linear function of the numbers of contraplex-diplex transitions involved in the system of dynamic equilibria of the electromers of benzene.

E. The Origin of Each Absorption Band.

One other question remains to be considered; namely, that of the possibility of relating each of the seven absorption bands to its proper source. It has been assumed that the vibrations of two secondary contraplex-diplex transitions are synchronous with light waves of frequency 3725 (band One) while the vibrations of twenty-four, i.e., the eighteen secondary and the six primary contraplex-diplex transitions, are synchronous with light waves of frequency 4200 (band Seven). The intervening whole numbers, 4, 7, 11, 16, and 20, represent the numbers of transitions smilarly involved in the production of bands Two, Three, Four, Five, and Six, respectively. Of the seven distinct groups of secondary phasotropic equilibria, one and only one group of transitions is related to band One. These two transitions of one group plus two transitions of one group plus two transitions of one of the seven closurs gives four which are related to band One. These two transitions of one group plus two transitions of one other of the seven

four plus three other transitions gives seven which are related to band Three. In this manner the entire series of whole numbers.



2, 4, 7, 11, 16, 20, 24 may be built up, but each increment in the series must involve the addition of the transitions of one and only one of the seven groups of secondary contraplex-deplex equilibria,

white one or more of the primary groups may or may not be added. This condition preserves the correlation of the seven groups of secondary phasotropic equilibria with the seven bands in a consistent and symmetrical manner.

The following scheme embodies such an an angement of the several groups of transitions, and should be studied in conjunction with the tables of primary and secondary phasotropic equilibria and the complete system of the electromers of benzene in dynamic equilibria. The groups of secondary equilibria are designated by the Roman numerals I. to VII inclusive and the primary equilibria by the letters $a, \beta, \gamma,$ and δ , while the subscript numeral is the number of transitions involved in the given group. For example, (V_2) signifies the two secondary contraplex-diplex transitions, namely, $C \neq C \Rightarrow D \neq D'$ and $C \Rightarrow C \Rightarrow D' \Rightarrow D'$ and group $V_1 \in \{\beta\}$ signifies the one primary contraplex-diplex transition, namely, $C \Rightarrow C \Rightarrow C'$, of group β . The symmetrically evolved scheme relating each of the seven absorption bands to its possible source is as follows —

Bard	Ongui	Number of Transitions Involved
One Two Three I our I've Six Seven	$ \begin{array}{l} V_{ij} \\ V_{i$	

Note that the transitions of group IV, involve the centric electromers C and D which constitute the nucleus, so to speak, of the complete system of dynamic equilibria of the electromers of benzene. Accordingly, the other groups of transitions are successively and collectively embraced, producing in a natural sequence the series of seven whole numbers which are functions of the oscillation frequencies of the seven bands, and which represent the number of specifically indicated contraplex-diplex transitions involved in the production of each band (espectively.

The conception that contraplex-diplex transitions occasion the absorption of light in carbon compounds is extended, in the next two chapters, to the dynamic formulæ and the absorption

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spectra of chlorobenzene, homobenzene, and naphthalene, with the uniform result that the oscillation frequencies of the absorption bands are functions of the numbers of contraplex-diplyx transitions involved in the systems of dynamic equilibria of their electromers.

CHAPTER XIX.

ABSORPTION SPECTRA AND DYNAMIC ELECTRONIC FORMULÆ OF CHLORO, BROMO, AND 10DO-BENZENE,

THE purpose of the present chapter is to test the validity of the absorption theory (just developed in relation to benzene) by extending it to the ultraviolet absorption spectra of chloro- and biomobenzenes. In this connection, the non-selective absorption of iodobenzene requires an explanation

The existence of a rule has been indicated. This rule, a linear relationship (r = xr + b), involves definite numbers (s) of contraplex-diplex transitions occurring within the systems of phasotropic equilibra of the electromers of the compound, and the oscillation frequences (r) of its absorption bands. In this and subsequent chapters, contraplex-diplex transitions will be termed absorption transitions. In other words it will be maintained that the oscillation frequency (r) of the head of a given band is a linear function of the number of absorption transitions (x) which function as the origin of the band of frequency (r) the function as the origin of the band of frequency (r).

Purvis 1-20 undertook the investigation of the absorption spectra of chloro, bromo, and todobenzenes in order to assertain (1) the nature of the absorption of the radiant energy, and (2) how far the displacement of one atom of hydrogen of benzene by chlorine, bromine, or rodine, affects the type of absorption when the compounds were in the vapour state, in solution in alcohol, or in very thin films. Therefore, the present chapter is limited to a discussion of the effect of the displacement of one hydrogen atom of the benzene molecule, by the halogens noted above, upon the absorption spectra of the compounds in solution (a) in alcohol, and (b) in thin films. The purpose of the present development is threefold.

- (1) To compare the data of Purvis with the data of Baly concerning the oscillation frequencies of the absorption bands of chlorobenzene when dissolved in alcohol.
- (2) To compare and interpret the differences in the values of the oscillation frequencies (according to Purvis) of the absorption

bands of chlorobenzene and bromobenzene under different physical conditions, namely (a) when the substances are dissolved in alcohol, and (b) when the substances are in the form of vy y thin films

(3) To present a new explanation of the non-selective absorp-

Purvis 121 found that alcoholic solutions of chlorobenzene and biomobenzene of equimolai concentrations, through equal thicknesses, exhibit seven wide diffuse bands which are comparable in appearance, and are shifted a little more towards the less refrangible end of the spectrum in the biomobenzene solution. Purvis thus demonstrated that the earlier investigations of Baly and Collie.125 and Balv and Ewbank,126 were incomplete because they failed to locate and determine the positions of the entire seven bands of these compounds Subsequently, Baly 127 again undertook the investigation of the absorption spectra of chlorobenzene and found seven bands, the same number previously established by Purvis. The positions of the heads of the bands, as determined by Purvis in wave-length values (a), have been converted in the present discussion to the corresponding values in oscillation frequencies (1/A) so that comparisons may be made readily with the data of Balv, originally given in oscillation frequencies. Furthermore, it is an advantage to have all data recorded in oscillation frequencies since the rule under consideration states that the oscillation frequencies are linear functions of the absorption transitions.

The following Table I. includes the oscillation frequencies of the absorption bands of benzene, chlorobenzene, and bromobenzene in alcoholic solution; and chlorobenzene and bromobenzene in very thin films. Iodobenzene shows no absorption hands.

TARLE I

Number	CgH ₅ H	C _b H ₅ Ci	CuH5 , Ci	GaH ₅ Be	CuHa Cl	C _G H ₅ . Br
of	Solution	Solution	Solution	Solution	in Films	In Films
Band.	(Baly)	(Baly)	(Purcis)	(Purvis)	(Purvis).	(Purvis)
One	3725	3684	3685	3679	3674	3670
Two	3765	3777	3781	3775	3772	3768
Three	3830	3825	3814	3821	3818	3815
Four	3915	3878	3880	3874	3871	3868
Five	4025	3920	3923	3977	3912	3909
Six	4110	3975	3984	3976	3974	3971
Seven	4200	4072	4082	4073	4054	4049
Jeven	4200	40/2	4002	40/3	4054	4

An inspection of the above table shows that chlorobenzene and bromobenzene either in alcoholic solution or in thin films deplay the same number of absorption bands as does benzene, namely, seven. Furthermore, the regions of selective absorption these compounds are not widely different. These facts lead to the conclusion that the absorption transitions which constitute the origin of the seven absorption bands of benzene must function similarly as the origin of the seven absorption bands of chlorobenzene and of bromobenzene. Therefore, before considering in detail the origin of the absorption bands of chlorobenzene and bromobenzene, it will be necessary to recall briefly the relationship between the oscillation frequencies of the seven bands of benzene and the numbers of absorption transitions which are presented by its electromers in dynamic equilibra.

Within the complete scheme of dynamic equilibria of the electromens of benzene (see Fig. 1, ρ . 188) there are involved altogether twenty primary systems and twenty-eight secondary systems of phasotropic equilibria. All of these transitions are tabulated on page 194, but it is more convenient in the present chapter to refer to the following abbievated Table II. which embodies only those transitions which function in the absorption of light, namely, the contraplex-diplex or absorption transitions. These are indicated as groups I to VII. and groups α , β , γ , and δ

A mathematical relationship between various groups or numbers of these absorption transitions and the oscillation frequencies of the seven absorption bands of benzene was developed by means of the system of rectangular co-ordinates in section D of Chapter XVIII.

Before extending this method of deriving the series of

numbers of absorption transitions to chlorobenzene and bromobenzene, it should be observed that each of these compounds is a mono-substituted derivative of benzene and the substituents. chlorine and bromine, are saturated atoms possessing neither free nor latent valences which might interfere in some manner with the centric valences of the benzene nucleus. Therefore, it is reasonable to assume further that the benzene nucleus in chlorobenzene and in bromobenzene is capable of undergoing the same centric-rearrangements 128 that are characteristic of benzene Accordingly, chlorobenzene and biomobenzene would likewise present within the scheme of dynamic equilibria of their electromers the same total number of absorption transitions that are common to benzene, namely, twenty-four. Hence the scheme of transitions for benzene indicated in Table II. (p. 204) and Fig 1 (p. 188) will apply equally to chlorobenzene and bromobenzene in the following discussion.

A. The Absorption Spectrum of Chlorobenzene in Alcoholic Solution.

In the following Table III, the oscillation frequencies of the absorption bands of chlorobenzene in alcoholic solution (as separately determined by Baly and by Purvis) and their respective successive differences are indicated. There is also included for subsequent reference a third column containing the frequencies of the bands of bromobenzene and their successive differences, the observations being made upon the compound in alcoholic solution -

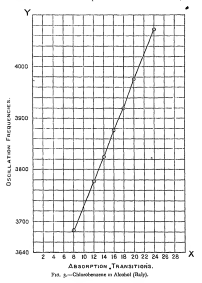
TABLE III

Number of Band.	CaHa Cl (Baly)	Difference	C _i H ₅ C1 (Purvis)	Difference.	C ₆ H ₅ Br (Purvit)	Difference
One .	3682	95	3685	96	3679	96
Two	3777	48	3781	33	3775	46
Three .	3825	53	3814 .	66	3821	53
Four Five	3878	42	3880	43	3874	43
Six .	3975	55	3923 3984	61	39 ¹ 7 39 ⁷ 6	59
Seven .	4072	97	,082	98	4073	97

206

The columns of differences show that the greatest discepanies occur in the frequencies of bands Thee, Six, and Seven. These differences are of such interest as to warrant a critical companison and discussion. In order to compare the data of Puvis with the data of Buly, some bass for comparson must be selected. A basis of comparison presents itself in the series of integral numbers of absorption transitions which are functions of the frequencies of the several absorption bands of chlorobenzene This series of numbers for chlorobenzene (in alcoholic solution) may be determined by employing the system of rectangular co-ordinates according to the principles and method previously described in the study of the absorption spectrum and dynamic formula of benzene.

First, the application will be made to Balv's data on the absorption spectra of chlorobenzene in alcoholic solution the following Fig. 3, the frequencies are indicated on the Y-axis and the absorption transitions on the X-axis. According to the present hypothesis the entire 24 absorption transitions function as the outsin of the band of highest frequency, namely, 4072; hence the point (4072, 24) Now there can be found passing through this point only one straight line which will intersect the abscissæ from the frequency values (Y-axis) at points which have corresponding values on the X-axis (absorption transitions) approximately equal to whole numbers This straight line passes through the points (4072, 24) and (3682, 8) involving the bands of highest and lowest frequencies respectively, and the series of whole numbers of absorption transitions which are the linear functions of the frequencies of the seven bands of chlorobenzene are thus found to be 8, 12, 14, 16, 18, 20, 24. The line which relates these numbers with the corresponding frequencies is an approximately straight line. It is important to note that according to the present hypothesis this line should be absolutely straight provided that the data for the oscillation frequencies as determined by Baly are absolutely correct. Be this as it may, the deviations from the straight line are so slight that the series of whole numbers thus ascertained will here be provisionally accepted and employed for purposes of comparison in their relation to Baly's data for chlorobenzene in alcoholic solution. On the other hand, if the values for the oscillation ficquencies as determined by Purvis be as nearly correct as the data of Baly, they should be related likewise to the same series of whole numbers of absorption transitions. Therefore, an



application of the hypothesis to the data of Purvis naturally follows.

In the following Fig. 4 the frequencies of the absorption bands of chlorobenzene in alcoholic solution as determined by Purvis are indicated on the Y-axis; the absorption transitions, on the X-axis. Band Seven of highest frequency, 4082, involving 24 absorption transitions, establishes the point (4082, 24).

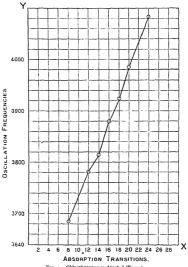


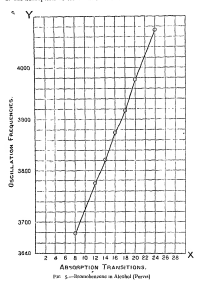
Fig. 4.-Chlorobenzene in Alcohol (Purvis)

Now, in this instance there cannot be found any straight line passing through this point which will intersect the abscissæ from the frequency values (Y-axis) at points which have correspond-

ing values on the X-axis (absorption transitions) approximately equal to whole numbers Furthermore, if the series of whole numbers related to Balv's data be applied to the data of Purvis. the line which relates these whole numbers with the corresponding frequencies is not a straight line (see Fig. 4) Therefore, in terms of the present hypothesis, the values for the frequencies of the absorption bands of chlorobenzene in alcoholic solution as determined by Purvis are not as accurate as the corresponding values determined by Baly Further evidence of the partial inaccuracy of the data of Purvis on chlorobenzene becomes evident when his data on bromobenzene in alcoholic solution are compared with Baly's data on chlorobenzene.

B. The Absorption Spectrum of Bromobenzene in Alcoholic Solution.

Again, if reference be made to Table III., it will be observed that there is a remarkably close agreement between the values for the oscillation frequencies of the absorption bands of chlorobenzene as determined by Baly and the corresponding frequencies of the bands of bromobenzene as determined by Purvis This naturally leads to the assumption that the series of whole numbers (8, 12, 14, 16, 18, 20, 24) of absorption transitions which are linear functions of the frequencies of the bands of chlorobenzene are likewise linear functions of the frequencies of the bands of bromobenzene. That such is actually the case is shown by again employing the system of rectangular co-ordinates. Fig. 5 shows how closely the series of numbers of absorption transitions (X-axis) lend themselves to the formation of a straight line when plotted with the corresponding frequencies (Y-axis) of the bands of bromobenzene. The data of Puivis are employed in Fig 5 In other words, the present hypothesis shows that the frequencies of the absorption bands of chlorobenzene, as determined by Baly, and also those of bromobenzene as determined by Purvis, are linear functions of the same series of numbers of absorption transitions. This is not quite the case with the frequencies of the hands of chlorobenzene as determined by Purvis and discussed in the preceding Section A. Therefore, they cannot be as commensurately exact as the corresponding determinations of Balv. A more definite, in fact, a ugid mathematical comparison of all of the data under consideration will be presented as soon as the values for, the oscillation frequencies of the absorption bands of chlorobenzene and bromobenzene in



thin films have been considered. All of the data under consideration will then be subjected to an application of the method of least squares.

C. Absorption Spectra of Chlorobenzene and Bromobenzene in Thin Films.

Purvis has found that very thin films of these two substances exhibit seven wide diffuse bands which are comparable in appearance and resemble the solution bands, but they are shifted more towards the less refrangible regions. The bands of the bromohenzene films are shifted more towards the less refrangible region than those of the chlorobenzene films. The frequencies of the bands of these compounds and their respective successive differences are presented in the following Table IV

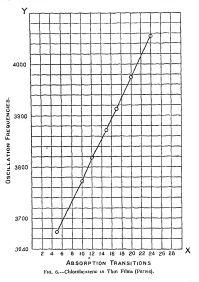
TABLE IV.

No of	Band	C _{fi} ll ₅ Cl. (1hm Filma)	Difference	CgHs, Br (Thm Filmss)	Difference
One Two Three Four Five Six .		367‡ 3772 3818 3871 3912	98 46 53 41 62	3670 3768 3815 3868 3909	98 47 53 41 64
Seven		4054		4049	

The columns of differences clearly show that the relative positions of the heads of the bands of each of these compounds are identical. Accordingly, the frequencies of the bands of chlorobenzene and of bromobenzene in thin films likewise should be related to the same series of numbers of absorption transitions. This series, however, cannot be the same as that for the bands of chlorobenzene and bromobenzene in alcoholic solution because the relative positions of bands Six and Seven of the compounds in alcoholic solution are quite different from the relative positions of the same bands of these compounds in thin films. The maximum difference between the sixth and seventh bands in the former case is 98, in the latter case, 80

The series of numbers of absorption transitions which are linear functions of the frequencies of the bands of chlorobenzene and of bromobenzene when in thin films may be determined

according to the method previously described for these compounds when in alcoholic solution. In the following Fig. 6 the frequencies of the bands of chlorobenzene are indicated on



the Y-axis; the absorption transitions, on the X-axis. Band Seven, frequency 4054, involving 24 transitions, establishes the point (4054, 24) through which only one straight line may be

found to pass which will intersect the abscisse from the frequency values (Y-axis) at points which have corresponding values on the X-axis (absorption transitions) equal to whole numbers. This line establishes the series of transitions (5, 10, 12, 15, 17, 20, 21) which are functions of the frequencies of the corresponding bands (One to Seven) of chlorobenzene in thin films. The same series of whole numbers also functions for the frequencies of the bands of bromobenzene in thin films because the relative positions of the heads of the bands of chlorobenzene and bromobenzene are identical.

D. Application of the Method of Least Squares to Absorption Data.

A more rigid mathematical comparison of all of the data under consideration in this paper is helewith presented in the following Tables V. to IX. inclusive,

TABLE V. CHLOROBENZENE (in alcohol) y = 24.41661 + 3484 gos

A	B (Baly)	С	D
8 12 14 16 18 20 24	3682 3777 3825 3878 3920 3975 4072	3680 237 3777 904 3826 737 3875'570 3974'103 3973'237 4070 903	+ 1 763 - 0'904 - 1'737 + 2'430 - 4 4'93 + 1'763 + 1'097

TABLE VI Chlorobenzene (in alcohol) y = 24 9806x + 3478.738.

A	13 (Purvis)	C	D
8	3685	3678 583	+ 6417
12	3781	3778 505	+ 2 49
I t	3814	3828*466	- T4'466
10	3880	3878 428	+ x 572
18	3923	3928 389	- 5389
20	3984	3978 350	+ 5 650
24	4082	4078 272	+ 3 728

TARLE VIE

Bu	OMOB	NZFNI	(in a)re	(lode	7 = 2	1 60011	1- :	34/5.2	-24	
-					-	_			-	
A		В (Pussi)	-	L			D	è	
				-						

A	B (Pust)	Ĺ	D
5	3679	3676 017	. ⊢ ≥ 953
14	3775	3771 809	10191
rı	382r	3821.180	- 3.180
16	3874	3873*570	+ 0.430
18	3917	39.22"751	- 5 95 ¹
20	3976	3972 332	4 3 668
21	4073	4071 093	+ 1.907

TABLE VIII CHI ORORENZENE (in thin films) p = 19 97171 + 3573 986.

Α	B (Purer)	C	D
		~	
5	3674	3673 X14	+ 0.120
10	3774	3773 705	- I'705
12	3818	3813 646	+ 4 351
15	3871	3573 561	- 2 561
17	3012	3913 505	- 1'505
20	3974	3973*420	+ 0.290
24	1051	10521207	al 0 503

TABLE IX. Brostopenzi ne (in thui films) v = 10.9548v + 3570.665

B (Pursis)	С	D	
3670	3570.130	- 0 430	
3768	3770 213	- 2,413	
3815	3810'122	+ + 878	
3868	386g g87	- 1'087	
3909	3909 896	~ o 8g6	
397I	3969 76t	+ 1.530	
1049	4019 580	- 0.280	
	3670 3768 3815 3868 3909 3971	3670 3670°139 3768 3770°213 3815 3810°122 3868 3869 987 3909 3909 396 3971 3969 761	

In each of the above Tables (V.-IX.), column A contains the theoretically determined series of whole numbers of absorption transitions which function as the origin of the corresponding oscillation frequencies respectively indicated in column B. The equation for the straight line for each table of data expresses the relationship between the number of absorption transitions (x) and the corresponding oscillation frequency (y). In each case the equation was derived by applying the method of least squares to the theoretically determined numbers of absorption transitions (x) in column A, and the experimentally determined frequencies (y) in column B. By means of these equations, values for the oscillation frequencies have been calculated and are indicated in column C. The deviations of the calculated from the experimentally determined frequencies (i.e., B-C) are recorded in column D

It will be observed that, with a few exceptions, the deviations show a remarkably close agreement between the calculated and the observed frequencies. Furthermore, a critical test, both of the validity of the rule that the frequencies are linear functions of the numbers of absorption transitions, and of the accuracy of the data in question, may be found by calculating the values for the probable error by means of the application of Peters' formula for each of the columns of deviations in the above Tables V.-IX. Of course, it is understood that the term "probable error" does not mean that said error is more probable than any other. It signifies that in any subsequent observations the probability of committing an error greater than the probable error is equal to the probability of committing an error less than the probable error. Therefore, the determination of the probable error for each of the above tables of data will give a numerical value which will represent the relative degree of accuracy of the several sets of data in the above tables, and also serve as a test of the validity of the rule. The following tabulation comprises the results of the application of Peters' formula for the probable error of a single observation -

 $r = 0.8453 \frac{\Sigma v}{\sqrt{n(n-1)}}$, $\Sigma v = \text{sum of the deviations for each set}$ of data, n = number of measurements of frequencies, namely 7,in each table of data.

Tables.		2,0	r.
V	Chlorobenzene in alcohol (Haly) Bromobenzene ", ", {Purvs} Chlorobenzene ", tim hins (",) Bromobenzene ", ", " (",)	14'0970	1.838
VI		39 7169	5.179
VII		18'2891	2.385
VIII		11 5537	1.507
IX,		12 2316	1.595

A comparison of the values in the above table shows that the probable error of a single observation for each table of data (with the exception of VI. and possibly VII., i.e., chlorobenzene and bromobenzene in alcohol according to Purvis), is practically within the limits of error of experimental observation.

The developments presented in this chapter warrant the following conclusions —

- (1) The oscillation frequencies of the absorption bands of chlorobenzene and of bromobenzene, either in alcoholic solution or in thin films, are linear functions of a corresponding series of whole numbers. In terms of the present hypothesis, the whole number which is related to a given frequency represents the number of absorption transitions which function as the origin of the band of given frequency.
- (2) One series of numbers of absorption transitions applies equally to the data for chlorobenzene (Baly) and bromobenzene (Puivis) when observations were made of the compounds in alcoholic solution.
- (3) Another series of numbers of absorption transitions applies equally to the data of Purvis for chlorobenzene and for bromobenzene when observations were made of the pure substances in thin films.
- (4) The probable errors for the observations of the substances in thin films are not only more nearly equal but they are also smaller than the corresponding probable errors for the observations of the substances in alcoholic solution.
- (5) The possibility of representing the frequencies of the absorption bands of a compound as linear functions of a series of whole numbers affords a means of determining the relative accuracy of the observations. For example, the data of Purus for chlorobenzene in alcoholic solution (Table VI.) do not conform to this condition as is evident in Fig. 4. Furthermore, the probable error for this set of observations is not commensurate with the probable errors for the other tables of data, and therefore indicates greater errors in the determinations of the frequencies of the absorption bands of chlorobenzene than in the other data under discussion."

E. The Origin of the Absorption Bands.

There now remains for consideration the possibility of 1elating each of the seven absorption bands of chlorobenzene and bromobenzene to its probable source. In other words, in terms of the present hypothesis each absorption band owes its origin to the existence of certain numbers and groups of absorption transitions. The groups of absorption transitions which are common to benzene, chlorobenzene, and bromobenzene have been indicated in Table II. The principles employed in relating the seven bands of benzene to their respective groups of absorption transitions have been described in Chapter XVIII., Section E. They are applied, herwith, in the same manner to the absorption bands, of chlorobenzene and bromobenzene.

Two different series of numbers of absorption transitions have been indicated in this chapter, namely, the series (8, 12, 14, 16, 18, 20, 24) for either chlorobenzene or bromobenzene in alcoholic solution, and the series (5, 10, 12, 15, 17, 20, 24) for either chlorobenzene or bromobenzene in thin films. Accordingly, the two different schemes presented in the following Tables X. and XI. embody the possible airangements of the several groups of absorption transitions which may function as

TABLE X.
CHLOROBENZENS AND BROMOBENZENE IN ALCOHOLIC SOLUTION,

Band	Origin.	Number of Absorption Transitions.
One Two Three Four Five Six Seven	$\begin{split} & \frac{1 V_2 + \alpha_0 + \beta_1 + \gamma_1 + \beta_2}{1 V_2 + \alpha_0 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VII}_3]} & VII_3 \right) \\ & VI_2 + \alpha_0 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VII}_3] \\ & VI_2 + \alpha_0 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VII}_3] + [II_2 \text{ or VI}_2] \\ & VI_2 + \alpha_1 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VII}_3] + [II_2 \text{ or VI}_2] + [III_2 \text{ or VI}_3] \\ & VI_2 + \alpha_2 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VI}_3] + [II_1 \text{ or VI}_2] + [III_2 \text{ or VI}_3] \\ & VI_2 + \alpha_2 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VI}_3] + [I_1 \text{ or VI}_4] + [II_1 + V_2] \\ & VI_2 + \alpha_2 + \beta_1 + \gamma_1 + \beta_2 + [I_1 \text{ or VI}_4] + [I_1 + V_1 + III_2 + V_2] \end{split}$	8 12 14 16 18 20 24

TABLE XL.

CHLOROBENZENE AND BROMOBENZENE IN THIN FILMS.

Band	Origin	Number of Absorption Transitions
Three	$\begin{split} & V_{+} + \{(a_{2} + \beta_{3}) \text{ or } (\gamma_{1} + b_{3})\} \\ & V_{+} + \{(a_{2} + \beta_{3}) \text{ or } (\gamma_{1} + b_{3}) + (\beta_{3} \text{ or } \gamma_{1}) + (I_{3} \text{ or } VII_{3}) + (I_{4} \text{ or } VII_{3}) \\ & V_{+} + \{(a_{3} + \beta_{3}) \text{ or } (\gamma_{1} + b_{3}) + (\beta_{3} \text{ or } \gamma_{1}) + (I_{3} \text{ or } VII_{3}) + (II_{3} \text{ or } VI_{3}) \\ & V_{+} + a_{3} + \beta_{3} + (\beta_{3} \text{ or } \gamma_{1}) + (I_{3} \text{ or } VI_{3}) + (II_{3} \text{ or } VI_{3}) \\ & V_{+} + a_{3} + \beta_{3} + (\beta_{3} + \gamma_{1}) + (I_{3} \text{ or } VI_{3}) + (II_{3} \text{ or } VI_{3}) \\ & V_{+} + a_{3} + \beta_{3} + (\beta_{3} + \gamma_{1}) + (I_{3} \text{ or } VI_{3}) + (II_{3} + VI_{3}) \\ & V_{+} + a_{3} + \beta_{3} + (\gamma_{3} + I_{3} + VI_{3}) + (II_{3} + VI_{3}) \\ & V_{+} + a_{3} + \beta_{3} + (\gamma_{3} + I_{3} + VI_{3}) + (II_{3} + VI_{3}) \\ \end{split}$	5 10 12 15 17 20 24

the origin of the corresponding absorption bands. The notation is the same as that previously employed. For example, the symbol IV, represents the two absorption transitions of group IV., namely, $C' \Rightarrow C \Rightarrow D \Rightarrow D'$ and $C'' \Rightarrow C \Rightarrow D \Rightarrow D''$.

In comparing the above schemes it will be observed that the absorption transitions of group IV, involve the centuc electromers C and D (see Fig 1, p. 188) which constitute the nucleus, so to speak, of the complete system of dynamic equilibria of the electromers of benzene, chlorobenzene, and bromobenzene. The other groups of transitions are successively and collectively embraced, producing in a natural sequence the series of seven whole numbers which are linear functions of the oscillation frequencies of the seven bands, and which may probably represent the number of the specifically indicated absorption transitions involved in the production of the corresponding absorption band

It has also been noted that the series of numbers of transitions related to the absorption bands of chlorobenzene and biomobenzene in alcoholic solution is not identical with the series of numbers of transitions related to the absorption bands of these compounds in thin films The explanation of this difference may be found in the fact that in alcoholic solution the molecules of the dissolved compound cannot be as closely compacted as they are in thin films of the pure substance. Consequently, the relative positions of the electromers of the compound are different and this in turn may lead to different arrangements of the various groups of the electromers so that one series of groups of transitions would function as the origins of the bands of the compound in solution while another series of groups of absorption transitions would determine the origins of the bands of the compound in the oure state, that is, in thin films,

F. The Non-Selective Absorption of Iodobenzene.

The absorption spectra of solutions of rodobenzene have been investigated by Pauer 129 who found no bands Purvis 130 also studied the absorption spectra of various concentrations of alcoholic solutions of iodobenzene, and of thin films of the pure substance No bands were found in either case, whereas each of the corresponding mono-substituted derivatives of benzene. chlorobenzene and bromobenzene, exhibited seven bands. This anomalous behaviour on the part of iodobenzene demands an explanation

Puvis has offered the explanation that "the heavy iodine atom is the controlling force, and it damps and dislocates the movements of the atoms of the benzene nucleus as well as the alkyl side chains, so that the hythmical oscillations or vibrations are destroyed, and no selective absorption is possible". It is unidoubtedly true that the mass, the intrinsic characteristics, the orientation of the atoms of the benzene nucleus and its substituents, and the physical conditions of the vibrating system, may all function in determining the nature of the absorption spectra, but the absorption hypothesis of Puvis and his explanation of the non-selective absorption of nodobenzene and its derivatives must be recarded as deficient for the following reasons:

- (I) Purvis fails to define the nature or type of the rhythmical oscillations or vibrations of the so-called "oscillation centres" 141
- (2) The assumption that the weight of the iodine atom in iodobenzene damps and dislocates the movements of the atoms of the benzene nucleus, thereby preventing selective absorption, is somewhat arbitrary in that it fails to take into consideration another equally probable condition, namely, that the weights of other atoms, chlorine and bromme, which replace one hydrogen atom of benzene likewise may damp, or at least dislocate, the movements of the atoms of the benzene nucleus and, thereby, either after or prevent selective absorption. This, however, is not the case since chlorobenzene and bromobenzene each show seven absorption bands—the same number exhibited by bonzene. An inspection of the following tabulation further emphasizes the significance of this criticism:—

A		В	С	
C ₆ H ₅ . H C ₆ H ₅ . Cl C ₆ H ₅ . Br C ₆ H ₅ . I		H = 1 008 Cl = 35'46 Br = 79'92 I = 126'92	35'46 1'008 = 35'17 79'92 : 35 46 = 2 25 126 92 : 79'92 = 1'58	

Column A contains the formulæ of the compounds under consideration. Column B indicates the atomic weights of the substituents which replace one hydrogen atom of benzene; benzene, in turn, beng regarded as phenyl hydride, a monosubstituted derivative. Column C embodies data showing that

the chlorine atom which replaces one hydrogen atom is 35.17 times as heavy as the replaced hydrogen atom of benzene. Notwithstanding this very great difference in the ratio of the weights of the substituents chloring and hydrogen, chlorobenzene shows the same number of absorption bands as does phenyl hydride or benzene Furthermore, the bromme atom which may be regarded as replacing the chlorine atom is 2'25 times as heavy as the displaced chloune atom, nevertheless, the resulting compound, bromobenzene, shows seven absorption bands, Finally, the iodine atom replacing the bromine atom is only 1.58 times as heavy as the displaced bromine atom, but iodobenzene shows no absorption bands. Therefore, in view of these decreasing ratios of the weights of the substituents, hydrogen, chlorine, bromine, and iodine, to one another, the nonselective absorption of rodobenzene should not be attributed to the weight of the fodine atom. How then is the non-selective absorption of iodobenzene to be explained?

The present hypothesis has explained selective absorption by the occurrence of definitely described rearrangements of valencies within the electronic formule of the compound in dynamic equilibria. These rearrangements, contraplex-diplex transitions of absorption transitions, must be interfered with in some definite way if the selective absorption of the compound is to be prevented. Therefore, if chlorobenzene and bromobenzene each shows seven absorption bands and jodobenzene shows none, it must be concluded that the chlorine and bromine atoms in chlorobenzene and bromobenzene do not prevent the occurrence of absorption transitions, but that the iodine atom in iodobenzene (or its derivatives) inhibits, in some manner, the occurrence of absorption transitions. Now the existence of absorption transitions has been shown to depend upon rearrangements of the centric valences of the nucleus. Accordingly, the non-selective absorption of rodobenzene must be due to the inhibition or prevention of the centric rearrangements of the benzene nuclei by the substituted rodine atom

Why does not the substituted chlorine and bromme atom in chlorobenzene and bromobenzene inhibit centic rearrangements and thereby prevent the occurrence of the absorption transitions and the consequent selective absorption of these compounds? The answer to this question may be found in the particular chemical nature of the substituents chlorine, bromine, and iodine, In chlorobenzene and bromobenzene the substituents are completely saturated, i.e., they form no addition compounds through the intermediate agency of the chloring or broming atoms. "On the other hand, the jodine atom in jodobenzene is unsaturated since it combines directly with chloring to form iodobenzene dichloride according to the equation,132

The existence of the compounds, iodosobenzene, C,H,IO, and iodoxybenzene, C.H.IO., affords additional evidence of the unsaturated condition of the iodine atom in iodobenzene. The corresponding derivatives of chlorobenzene and bromobenzene are unknown.

The manner in which the unsaturated iodine atom inhibits the centure rearrangements of the benzene nucleus may be made evident by considering first the structural formulæ of the compounds todobenzene dichloride, todosobenzene, and iodoxybenzene, namely,

In iodobenzene, iodine is univalent, in iodobenzene dichloride and iodosobenzene, tervalent; in iodoxybenzene, quinquevalent. In periodic acid, HIO,, structurally represented

$$H-O-I = O$$
, the maximum valence of iodine is seven. Hence

it may be concluded that the unsaturated iodine atom in iodobenzene may possess two, four, and possibly six free or potential valences which may be represented as follows:-

$$C_0H_0-I = C_0H_0-I = C_0H_0-I$$

This property of certain atoms to display a capacity for increasing their degree of saturation is well known, and generally the additional valences manifest themselves, or are called into play, so to speak, in pairs. Furthermore, the two valences of such a pair are of opposite sign or polarity. For instance, in the reaction, NH₃ + HCl → NH₄Cl, the nitrogen atom changes its valence from three to five, that is, the valence is increased by two. In terms of the electronic conception of positive and negative valences this reaction is represented as follows.—

$$NH_3 = NII_3 - + H$$
 $NH_3 + H - CI = H_3N$

Analogously the iodine atom in rodobenzene increases its valence by two when combining with chlorine, thus:—

Analogously the several possible degrees of saturation of the indune atom in iodobenzene may be represented as follows, the additional valences appearing in pairs —

In the last formula the iodine atom displays its maximum valence of seven. One valence unites the atom to the benzene ring, and each of the three pairs of latent or potential valences comprises one positive and one negative valence.

Of the six centric valences of the benzene nucleus, three are positive and three are negative. Hence, the centric valences may be regarded as consisting of three pairs of free or potential valences, each pair consisting of one positive and one negative valence. Therefore, all of the free or potential valences of job benzene may be indicated in the following structural formula .—



Now the unsaturated valences of the iodine atom and the centric valences of the benzene nucleus would most naturally engage or neutralize one another as may be indicated in the following formula in which the substituted iodine atom has been placed in the centre for the sake of symmetry —

This electronic formula definitely illustrates how the centric valences of the benzene nucleus may be either "bound" or interfeted with by the unsaturated valences of the substituted iodine atom. Therefore, the non-selective absorption of iodobenzene naturally may be attributed to the unsaturated state of the iodine atom, the free or potential valences of which bind or interfere with the centric valences of the benzene nucleus, thereby inhibiting centric rearrangements, and consequently preventing the occurrence of absorption transitions.

CHAPTER XX.

DYNAMIC ELECTRONIC FORMULÆ AND THE ULTRAVIOLET ABSORPTION SPECTRUM OF NAPHTHALENE,

THE electronic conception of valence and the absorption-transition hypothesis have been applied, respectively, to the constitution and to the ultravolet absorption spectra of benzene, chlorobenzene, and bromobenzene. They are extended, herewith, to the constitution and ultraviolet absorption spectrum of naphthalene.

A. Electronic Formulæ of Naphthalene.

Benzene nucleu of the centric type were composed of the five electronic types of carbon atom (see p. 49). Each nucleus embraced three pairs of the combined types I. and V, II. and IV, or III. and III., symmetrically co-ordinated. Only six centric electronic formulae, centric electroners of benzene, were possible. By extending this method of building up electronic formulæ to the constituent atoms of the naphthalene molecule, two and only two, perfectly symmetrical centric electromers are possible, namely, A and B of Fig. 7—

Note that electromer A is composed of carbon atoms of types II. and V.; B, of carbon atoms of types II., III., and IV., symmetrically co-ordinated The inclusion of carbon atom of type III with types II. and IV. in electromer B is noted specifically in the two carbon atoms that are not united to hydrogen atoms. In other words, these are the two carbon atoms common to the

two benzene rings which constitute the naphthalene molecule, It is very significant that the hydrogen atoms in positions 1, 3, 6, and 8 of the naphthalene ring are negative while those in positions 2, 4, 5, and 7 are positive.

It is quite possible to interpret many of the chemical properties and eactions of naphthalene in terms of the polarities of these hydrogen atoms of the electronic formula of naphthalene, somewhat as substitution in the benzene nucleus, and many chemical properties of benzene and its derivatives were interpreted fully in terms of the electronic formula of benzene. This monograph has limited its interpretation of chemical reactions chiefly to benzene and its derivatives. The electronic formula of condensed benzene nuclei, such as naphthalene, anthracene, and phenanthrene, are dealt with in relation to phenomena of light absoration and fluorescence.

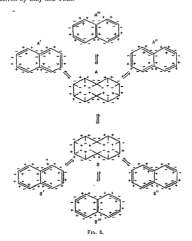
B. Systems of Dynamic Equilibria of the Electromers of Naphthalene.

A complete scheme of the systems of dynamic equilibia of the electromers of naphthalene is given in Fig. 8 (p. 226) The polarities of the valences which engage the hydrogen atoms of naphthalene are not indicated because they do not function in the centric rearrangements and related systems of equilibra.

Each electronic formula is to be regarded as the plane piopection of a space formula. The centric electromer A is the intermediate phase between the three possible phasotropic electromers A', A'', and A'''. By means of centric rearrangement, A, composed of carbon atoms of the types I and V., may be converted into B, composed of carbon atoms of the types II, III., and IV. In turn, B functions as the intermediate phase between the phasotropic electromers B', B'', and B''. This scheme presents six primary and nine secondary systems of phasotropic equilibria. Two of the primary and the nine secondary systems involve contraplex-diplex transitions which are assumed to constitute both the structural and the electronic explanation of the ultraviolet absorption spectrum of naphthalene.

Before tabulating and discussing the several systems of contraplex-diplex transitions, i.e., absorption transitions, it will be necessary to review briefly the relation between the absorption

spectrum of paphthalene and its chemical constitution as conceived by Baly and Tuck. 133



C. The Absorption Spectrum of Naphthalene.

Baly and Tuck state that "there are three absorption bands. namely, two narrow ones at 1/\(\lambda = 3125\) and 3220 respectively, and a broad band with its head at about $1/\lambda = 3700$ ". In attempting to correlate the position of these bands with the constitution of naphthalene Baly and Tuck maintain that "from the ease with which naphthalene is reduced in hot alcoholic solution by metallic sodium to the dihydro compound (I.),



and the further reduction to the tetrahydro compound (II), this being the final product of the reduction, there is little doubt that one of the rings is truly benzenoid, and the other contains two ethylenic double bonds, which according to Thiele's law, give their maximum effect at the two extremes; that is to say, at positions I and 4". Baly and Tuck then conclude that "Naphthalene therefore would seem to consist of two rings, of which one is truly benzenoid, and the other contains two conjugated double linkings. There is no reason to insist that the two rings are permanently endowed with one of the two above characters, in fact it seems that the interchange of these characters between the rings is perfectly possible, and no doubt is continually taking place."

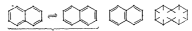
In seeking the origin of the three absorption bands in the spectrum of naphthalene, Baly and Tuck maist that the broad absorption band with its head at $1/\lambda=3700$ is due to the benzenoid motions of the naphthalene molecule and attibute this band to that half of the molecule which is benzenoid in character. On the other hand, they maintain that the two narrow bands at $1/\lambda=3125$ and 3220, which are nearer to the red end of the spectrum than any of the benzene bands, must be due to the isorropesis between the benzenoid tautomerism of the ring and the ethylenic double linkings of the other half of the molecule. Hence naphthalene is represented by the formula—

in which isorropesis between atoms 2 and 3 with the benzenoid system is indicated by the dotted lines

The foregoing suppositions of Baly and Tuck are open to the following three possible objections;—

(1) Since the naphthalene molecule is generally conceded to

be symmetrically constituted and to conform structurally to either of the three following symmetrical types:—



it naturally follows that any systems of vibrations of the naphthalene molecule should likewise be symmetrically developed and involve only symmetrical configurations

- (2) The fact that certain derivatives of naphthalene (such as the previously noted dihydro- and tetrahydro-naphthalenes) seem to consist of two rings, one of which is truly benzenoid while the other contains two conjugated double linkings, cannot be accepted as proof that naphthalene itself consists of two kinds of tings. Baly and Tuck partially admit this objection in their statement that "there is no reason to insist that the two rings are permanently endowed with one of the two above characters: in fact, it seems that the interchange of these characters between the rings is perfectly possible, and no doubt is continually taking place". This admission is incompatible with their fundamental assumption that one ring of the naphthalene molecule is benzenoid and that its vibrations produce band $I/\lambda = 3700$, while on the other hand the ethylenic ring permits of isorropesis between the atoms 2 and 3 with the benzenoid system, thereby accounting for bands 1/\(\lambda = 3125\) and 3220.
- (3) The hypotheses of Baly and others fail to indicate the existence of any quantitative relationship between the actual oscillation frequencies of the absorption bands and then proposed dynamic formula.
- In the proposed electronic formulæ for naphthalene and in the systems of dynamic equilibria of the various electromers there is perfect symmetry both in the structure of the electromers and in the transitions they undergo. Moreover, it is possible to show that a quantitative relationship in the nature of a linear function exists between the actual oscillation frequencies of the absorption bands of naphthalene and the numbers of absorption transitions which may be specifically indicated in the systems of dynamic equilibria of the various electromers.

D. Correlation of Oscillation Frequencies and Absorption Transitions.

Refering to the complete scheme of dynamic equilibria of the electromers of naphthalene (see Fig. 8), observe that there occurs only one centric rearrangement transition, namely, $A \simeq B$. The phasotropic electromers, A', A'', and A'', are each derived from the centric electromer A and each contains five dipler double bonds. They may therefore be regarded as mutually equivalent in these systems of equilibria in which they are involved. On the other hand, B', B'', and B''', each derived from the centric electroners B, are not mutually equivalent since B' and B'' each contain for contrapler double bonds while B''' contains four outraplers and one diplex double bonds B'' and B''' stands in a class by itself.* Thus there are three groups of phasotropic electromers, namely .—

 $A' = \bigcirc A'' = \bigcirc A'''$, each containing five diplex double bonds, $B' = \bigcirc B''$, each containing five contraplex double bonds, B'', containing one diplex and four contraplex double bonds,

The pumary and secondary systems of phasothopic equilibria are indicated in the following tables Those equilibria involving contraplex-diplex, or absorption, transitions are each followed by an asterisk:—



Only two of the primary systems (group a) present absorption transitions. The nine secondary systems are naturally divided into two groups. Group I, presents three transitions, each of which involves B" in dynamic equilibrium with A', A', and A'', respectively. Group II, presents six transitions

 ^{*} This type of an electromer functions in the production of the fluorescence band. Electrome formulæ in relation to fluorescence will be considered in the following chapter

involving B' and B" in dynamic equilibria with A', A", and A", respectively. Hence the following summary —

The correlation of the numbers of these transitions with the socillation frequencies of each of the bands in the ultraviolet absorption spectrum of naphthalene must now be developed. The oscillation frequencies of the heads of each of these bands, as determined by Balv and Tuck, are as follows:—

Band One			r/λ = 3125
Band Two			1/A = 3220
Band Three			I/A = 3700

Now the two transitions of group a and the three transitions of group I. (i.e., five transitions), involve the electromer B" which differs from each of the other phasotropic electromers of naphthalene. These five transitions may be assumed to function in the production of one of the absorption bands, presumably band One of lowest oscillation frequency, 3125. In other words, the vibrations of these five absorption transitions are assumed to be synchronous with light waves of frequency 3125. On the other hand, there remain the six absorption transitions of group II, which do not involve the electromer B" Hence the vibiations of these six absorption transitions may be synchronous with light waves of frequency 3220, i.e., band Two In other words, the oscillation frequencies 3125 and 3220 have been assumed to be functions of the whole numbers 5 and 6 respectively. The problem now demanding solution is the determination of that whole number which is a function of the oscillation frequency 3700 of the remaining band Three. How is this number to be derived?

Again, by employing the system of rectangular co-ordinates, possible to determine the number of absorption transitions involved in the production of band Three. In Fig. 9, the ftequencies are indicated on the Y-axis and the numbers of contraplex-duplex transitions on the X-axis. Five transitions have been assumed to function as the origin of band One of ftequency 3125, six transitions as the origin of band Two, frequency 3220. Now if a straight line be extended through the points (3125, 5) and (3120, 6) its extension will interest the

perpendicular from frequency 3700 at a p-int which has a corresponding value on the X-axis equal to the whole number eleven In other words, there are eleven absorption transitions functioning as the origin of band Three, frequency 3700. Furthermore, it is remarkable that this number 11, as thus derived, is equal to the sum of 5 plus 6. In terms of the absorption transition hypothesis this numerical result permits of only one conclusion, namely, that the five transitions which function as the origin of band One, and the six transitions which function as the origin of band Too, must function all together as the origin of the

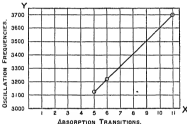


Fig. q.—Naphthalene (Baly and Tuck).

broad band Three. In other words, the vibrations of the entire II absorption transitions are synchronous with light waves of oscillation frequency 3700.

The scheme relating each of the absorption bands of naphthalene to its possible source is presented in tabular form as follows (the nomenclature is identical with that employed in the two preceding chapters):—

Hand		Or	igin			Number of Absorption Transitions Involved	
3126 3220 3700	$I_1 + \alpha_2$ II_6 $I_3 + \alpha_2 + II_6$:		5 6 11	

The equation for the straight line which relates the whole numbers, 5, 6, and 11, with the respective frequencies 3125 3220, and 3700, calculated according to the method of least squares, is (r = 0.68811 + 26.45013).

In the following table, column A contains the theoretically determined whole numbers represented by (x); column B, the experimentally determined oscillation frequencies represented by (p); column C, the frequencies as calculated from the equation y = 95.88711 + 264571613.—

A	В	C	D
 			-
5	3125 3220	312415968 3220 4839	+ 0 4032
11	3700	3699.0101	+ 0.0800

Note that the deviations, in column D, of the calculated from the actual oscillation frequencies are each less than one unit, a quantity exceedingly smaller than any deviation due to possible errors in experimental observation. These results further substantiate the hypothesis that the oscillation frequencies of the absorption bands of a given compound are functions of the number of absorption transitions involved in the systems of dynamic equilibria of its electromers.

The beating of these developments upon the question of colour and constitution is at once apparent since a coloured substance is one which exerts strong absorption within the ordinary limits of vision. Therefore the proposed absorption transition hypothesis which mathematically relates absorption and constitution should likewise function as the basis of the explanation of colour in relation to constitution.

CHAPTER XXI

FLUORESCENCE IN RELATION TO ELECTRONIC FORMULÆ.

SINCE it has been possible to interpret the absorption of light in terms of electronic formula: and absorption transitions, it naturally follows that fluorescence, notably manifested by such compounds as anthracene and phenanthrene, also may be interpreted by means of electronic formulæ and a new type of contraplex-diplex transition te mid "Buorescene transition".

Since fluorescence is produced only when the incident rays contain vibiations which the medium is capable of absorbing, it follows that the relation between fluorescence and absorption is to a certain extent reciprocal Not only can absorption of light cause fluorescence, but fluorescence in many cases, as shown by Burke, 184 increases the intensity of absorption. Hence, it is the purpose of the present chapter to extend the electronic conception of positive and negative valences and of contraplex-diplex transitions (as previously developed and illustrated in relation to absorption spectra in the three preceding chapters) to the interpretation of the phenomena of fluorescence. In other words, the relationship between chemical constitution and fluorescence will be considered from the standpoint of the existence of contraplexdiplex transitions within the systems of dynamic equilibria of the electromers of fluorescent compounds. A comprehension of the proposed fluorescence hypothesis necessitates, in the first place, a brief review of the foremost theories relating to fluorescence and constitution.

A. Fluorescence Theories in Relation to the Electronic Fluorescence Hypothesis.

A survey of fluorescent compounds by Rich Meyer 126 led to his "fluorophore" theory which, in its original form, serves as an excellent means of classifying fluorescent compounds, but it

affords no explanation of the relation between chemical constitution and fluorescence

As a result of the researches of J. Stark 126 on ultraviolet fluctescence, Meyer inverted his former view that the fluorophore is the seat of fluorescence, and both Stark and Meyer, 187 have concluded that in aromatic substances the bensene nucleus is the carrier of fluorescence, while the fluorophores and various substituents act so as to bring the fluorescent vibrations within the visible portion of the spectrum. The condensation of benzene nuclei accomplishes the same result

The extensive researches of Kauffmann 138 culminating in the lummophore and fluorogen theory also leads to the conclusion that in aromatic compounds the benzene nucleus is the seat of fluorescence, but this is not evoked until two kinds of groups -the auxochrome and fluorogen-have been introduced into certain positions. The introduction of the auxochrome excites luminescence, thereby indicating the approaching state of fluorescence. The subsequent addition of a luminophore perfects the process in the production of fluorescence.

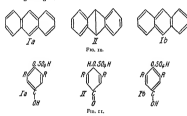
Francesconi and Bargellini 139 also admit the important part played in fluorescence by the benzene nucleus Recognizing that the observations of Meyer and Kauffmann are concerned chiefly with visible fluorescence, they claim that all aromatic compounds are fluorescent and that it is premature to attempt to explain the action of various substituents. Nevertheless, they classify substituents according to their action on the fluorescence of the parent substance. Those substituents which increase the fluorescence are called "auxoflores," and those which depress it, " bathoflores ".

It is important to the development of the electronic fluorescence hypothesis to note that the common point of agreement in each of the above theories is the tendency to relate the origin of fluorescence to the benzene nucleus.

The relation between tautomeric change and fluorescence was originally embodied in the theory of Wiedemann,140 that the molecule of a fluorescent substance exists in two forms, one of which is more stable than the other The stable form absorbs the energy of light vibration, and is thereby transformed into the less stable modification which spontaneously passes back to the stable form, emitting the previously absorbed energy as the

fluorescent light Wiedemann maintaine; that the two varieties of substance may be produced by the shifting of an atom within the molecule, and that the fluorescent light is directly due to the vibrations in the other which are set up by the motions of this atom.

Hewitt's theory "I" of double symmetric tautomerism involves the conception of Wiedemann that a fluorescent substance must exist in interchangeable forms. For instance, anthracene (Fig 10), and di-phenylpyrione-sulphate (Fig 11) present the following charges:—



In either of the above figures the molecule (Ia) passes to (II), and then to (Ib), whereupon the process is repeated in the reverse direction. The molecules (Ia) and (Ib) are chemically identical Hewitt likens these changes to the movements of a swinging pendulum The limiting positions in the amplitude of vibration correspond to forms (Ia) and (Ib) while the position of rest, so to speak, is represented by form (II).

It should here be noted that the changes from one form to another are accomplished in two distinct ways: (1) In anthracene the changes are due solely to a rearrangement of the positions of the double bonds, i.e., to changes in the direction of valences. (2) In di-phenylpyrone-sulphate, the rearrangement of double bonds is accompanied by a change in the position of a hydrogen atom. The significance of these changes from an electronic standpoint will appear in the definition of the term "fluorescence transition," which will be developed in a subsequent paragraph.

Let it now be recalled that Wiedemann maintained that fluorescence was due to vibrations set up by the motion or the wandering of an atom, but Drude 112 has shown that the vibrations of the atom itself, or the group of positive elections must correspond to the infra-red portion of the spectrum, while the periods in the visible and ultraviolet region are due to dispersional or valency elections. Therefore, the second form of the substance required by Wiedemann's theory need not be produced by the movement of an atom, but merely by the chance in position of an electron or valency. Since any change in the position of an atom is accompanied by a change in the positions of the double bonds. it follows that Hewitt's examples of double symmetric tautomerism given above will fulfil either one or both of these conditions

From this brief review of the foremost fluorescence theories. the following conclusions may be drawn :-

- (1) The origin of fluorescence in aromatic compounds is related m some way to the benzene nucleus
- (2) Physico-chemical evidence shows that the affinities of the benzene nucleus are in a state of continual oscillation
- (3) Physical theories require the presence of mobile negative electrons or valencies in the molecules of fluorescent compounds.

Therefore, any hypothesis which may be designed to express the relationship between chemical constitution and fluorescence must correlate these three conclusions

B. Fluorescence Transitions.

In the preceding chapters it has been shown that absorption of light is due to the existence of contraplex-diplex transitions within the systems of phasotropic equilibria of the various electromers of a given compound. These electronic systems of phasotropic equilibria also serve as examples of double symmetric tantomerism. Now note that any compound which contains a benzene nucleus (or condensed benzene nuclei) when considered from the standpoint of the electronic conception of positive and negative valence, will involve contraplex-diplex or absorption transitions within the primary and secondary systems of phasotropic equilibria of its electromers, and consequently must manifest one or more absorption bands in its spectrum. Now since fluorescence is produced only when the incident rays contain

vibrations which the medium is capable of absorbing, it naturally follows that the function of the benzene nucleus in fluorescent compounds is to make possible the absorption of light as the result of the existence of contraplex-diplex transitions. Thus the three conditions noted above are correlated, and an explanation of the factor of absorption in the phenomenon of fluorescence is their by afforded.

The various theories on fluorescence and chemical consistuation have failed to explain the factor of absorption in connection with the relations between the wave-lengths of the absorbed and the fluorescent or emitted light. It has been shown that incident light of a given wave-length may exerte a fluorescence consisting of several different rays; and, conversely, a given ray in the fluorescent spectrim may correspond to absorbed light of different wave-lengths. How are these facts to be interpreted in terms of contraplex-diplex transitions? In other words, what conditions in the making and breaking or rearrangement of contraplex and of diplex double bonds could be assumed to cause the emission of light of an oscillation frequency different from that which is absorbed? An answer to this question may be found in the following definitions of two possible types (I and II) of contraplex-diplex transitions.

Let A_1 and A_2 represent two electromers of a given compound in phasotropic equilibrium with one another. Also, let d_1 and d_2 be the number of diplex double bonds, and d_1 and d_2 be the number of contraplex double bonds in A_1 and A_2 respectively. Now if $d_1 = d_2$ and $d_1 = d_2$, then in the equilibrium $A_1 = d_2$ and the properties of the properties of light nor fluorescence. On the other hand, if d_1 is unequal to d_2 , or if d_3 is unequal to d_3 or if there would result two types (I. and II.) of contraplex-diplex transitions in the equilibrium $d_1 \Leftarrow d_2$.

Type I When $d_i = c_i$ and $c_i = \bar{d}_i$, the equilibrium $A_i \approx \lambda_i$ would involve contraplex-diplex transitions in which the number of diplex double bonds in one electromer is equal to the number of contraplex double bonds in the other electromer. In other words the number of diplex bonds in one electromer is balanced by an equal number of contraplex bonds in the other electromer. Such a condition, fully described and exemplified in preceding chapters, constitutes the origin of a absorption band, that is the absorption of light of a definite oscillation frequency synchronous with the rate of transition from the one electioner to the other. For convenience in reference, contraplex-diplex transitions of this type days been and will be termed "absorption transitions".

Type II. When $d_1 + c_2$ or $c_1 + d_3$, the equilibrium $A_1 \approx h_3$ will involve contraplex-diplex transitions in which either the number of diplex double bonds of the one electromer is unequal to the number of contraplex double bonds of the other electromer, or vice versa. Such a condition could be productive of two results simultaneously. (1) The absorption of light of a given frequency since some of the diplex or contraplex bonds of the one electromer are in equilibrium with, or are balanced by, an equal number of contraplex or diplex bonds, respectively, in the other electromer. (2) The existence of a residual number of unbalanced double bonds which do not function in the absorption of light, must function in the emission of light of a frequency different from that which is absorbed, hence fittorescence Contraplex-diplex transitions of this type (II.) will be termed "fluorescence transitions".

The several conditions described above may be summarized as follows:—

When $c_1=\delta_2$, and $d_1=d_2$, the transition $A_1\rightleftharpoons A_2$ causes neither absorption nor fluorescence,

When $e_1 + e_2$, or $d_1 + d_2$, two types (I and II) of contraplex-diplex transitions are possible.

Type I. When $d_1=c_2$ and $c_1=d_2$, transition $A_1 \rightleftharpoons A_2$ produces absorption only.

Type II. When $d_1 + c_2$, or $c_1 + d_2$, transition $A_1 \rightleftharpoons A_2$ occasions simultaneously, (1) absorption, and (2) emission of light; hence fluorescence

A general hypothesis may now be stated, namely, that a substance manifests fluorescence whenever within the systems of dynamic equilibria of its electromers the number of diplex dualite bonds undergoing rearrangement is unequal to the number of contraplex double bonds simulationally undergoing rearrangement.

From the method of its development it is evident that this hypothesis embedies the fundamental features of each of the presidently moted theories. It will now be extended to the dynamic formula and fluorescent spectra of anthracene and phenanthrene vapours.

C. Fluorescent Spectra of Anthracene and Phenanthrene.

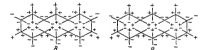
Elston ¹² has made an exhaustive study of the fluorescent and absorption spectra of anthracene and phenauthrene vapour and states that "the fluorescent spectrum of anthracene vapours consists of three bright bands at 300, 415, and 432 $\mu\mu$ superposed upon a continuous region extending from 365 to 470 $\mu\mu$. There is no evidence of lines . When the fluorescent spectrum of pure phenauthrene vapour was photographed, it was found to consist of the same bands as that of anthracene, but with an additional band at 360 $\mu\mu$ ". Elston then concludes that "there is an intimate connection between the fluorescence of the vapours of the two isometic substances, undoubtedly due to their common chemical composition (C₁₄H₁₀) and similar structural composition Just what gives use to the extra band in the fluorescent spectum of phenanthrene is not apparent "

Elston is undoubtedly correct in stating that there is an intimate connection between the fluorescence of the vapous a fint the two substances but this cannot be attributed entirely to their similar structural composition. While anthracene and phenanthiene each consists of three condensed benzene nuclei the position of the central nucleus in each formula genders them dissimilar. A glance at the relative positions of carbon atoms 9 and 10 in each of the following formulæ makes evident the difference in structure:—

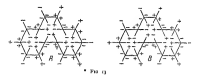
Anthracene and phenanthrene have three fluorescence bands in common. Elston states that the cause of the extra fluorescence band in the phenanthrene spectrum is not apparent. Now since these compounds are dissimilar in structure, it is natural to assume that the existence of the extra band is related in some way to this difference in structure. Hence in the application of the electionic conception of positive and negative valences to the constituent atoms of anthracene and phenanthrene, the purpose of the present chapter becomes threefold—

- (1) To show tifat within the systems of dynamic equilibria of the electromers of anthracene and phenanthrene there necessarily exist fluorescence transitions.
- •(2) To account for the existence of the extra band in the fluorescent spectrum of phenanthiene vapour
- (3) To show that the oscillation frequencies of the fluorescence bands of arthracine and plenanthrene may be represented as linear functions of the numbers of fluorescence transitions occurring within the respective systems of dynamic equilibria of their electromers.

The method of developing the electronic formulæ of benzene and naphthalene by symmetrically co-ordinating carbon atoms of types I. and V, II. and IV., and III. and III. has been applied to the constituent atoms of anthracene and phenanthrene. Two and only two centric electromers (A and B) of each of these compounds are derived. They are represented in Figs. 12 and 13 tespectively



F10, 12,



In each of the above figures the centric electromers (A) are composed of carbon atoms of the types I, and V while the centric electromers (B) embrace in their structure carbon atoms of the types II, III, and IV.

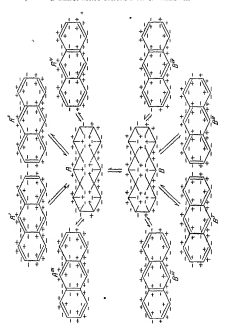
D. Systems of Dynamic Equilibria of the Electromers of Anthracene,

A complete scheme of the several systems of dynamic equilibria of the electromers of authorece is presented in Fig. 14.

Each electronic formula is the plane projection of a space formula. The centric electroner A is the intermediate phase between the four possible phasotropic electroners, A^{\dagger} , A^{\dagger} , A^{\dagger} , A^{\dagger} , and $A^{\dagger V}$. By means of centric rearrangement A is convertible into B. In turn, B functions as the intermediate phase between B^{\dagger} , B^{\dagger} , $B^{\dagger \dagger}$, B^{\dagger} , $B^{\dagger \dagger}$, and $B^{\dagger V}$.

The electromers of anthracene may be arranged into groups depending upon their respective symmetry and the number and kind of double bonds existing in their structure, thus:—

The primary and secondary systems of phasotropic equilibra which involve absorption and fluorescence transitions are included in the following table. (Note that the absorption transitions, such as $A^1 \Rightarrow A \Rightarrow B \Rightarrow B^1$ and $A^{111} \Rightarrow A \Rightarrow B \Rightarrow B^{112}$ are followed by a single asterisk. The fluorescence transitions, such as $A^1 \Rightarrow A \Rightarrow B \Rightarrow B^{111}$ and $B^{11} \Rightarrow B \Rightarrow B^{111}$ involving the electromers B^{111} and $B^{111} \Rightarrow B \Rightarrow B^{112}$ involving the electromers B^{111} and $B^{111} \Rightarrow B^{112} \Rightarrow B^{112} \Rightarrow B^{113} \Rightarrow B^{113}$



There are twelve fluorescence transitions within the complete system of dynamic equilibria of the electromers of anthracene.

E. Systems of Dynamic Equilibria of the Electromers of Phenanthrene.

A complete scheme of the several electromers of phenanthuene in dynamic equilibria is presented in Fig. 15.

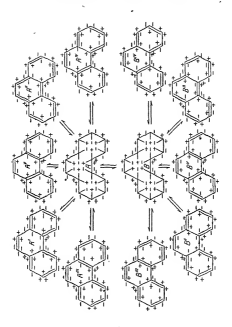
The centric electromer A is the intermediate phase between the five possible phasotropic electromers A^i , A^m , A^m , A^m , A^m , and A^v . By means of centric rearrangement A is convertible into B which functions as the intermediate phase between the electromers B^i , B^i

The electromers of phenanthrene may also be arranged into groups depending upon their respective symmetry and the number and kind of double bonds in their structure, thus:—

It should be observed that an electromer of the type B^V containing five contraplex and two diplex double bonds is not to be found among the electromes of anthracene. Hence the existence of A^V and B^V in the phenanthrene system serves to distinguish it from the anthracene system, and furthermore must bear some relation to the additional fluorescence band in the spectrum of phenanthrene vapour.

The primary and secondary systems of phasotropic equilibria involving absorption and fluorescence transitions are included in the following table.—

$$\begin{array}{c} \text{Primary Systems,} \\ \text{Primary Systems,} \\ \text{St.} = \text{But.} \\ \text{Br.} = \text{But.} \\ \text{Secondary Systems.} \\ \text{Secondary Sy$$



There are twenty-three fluorescence transitions within the state of dynamic equilibria of the electromers of phenanthrene, that is, eleven more than are to be found in the anthracene system. This additional number of eleven transitions is made possible thungth the existence of the electromers AV and BV.

The first purpose of this chapter, namely, to demonstrate the existence of fluorescence transitions within the systems of phasotiopic equilibria of the electromers of anthracene and phenanthrene, is realized in the preceding schemes of dynamic equilibria and tabulations thereof.

The second purpose, namely, to account for the existence of the extra band in the fluorescent spectrum of phenanthrene vapour, is realized in the existence of the electromers A' and B' of the phenanthrene system. Electromer B', containing 5 contralex and 2 diplex double bonds is a type which is not known in the authracene system. The existence of A', and B' makes possible eleven more fluorescence transitions in the phenanthrene system than in the authracene system. Hence these conditions, peculial to the phenanthrene system, may naturally be assumed to constitute the explanation of the additional fluorescence band in the phenanthrene spectrum.

F. The Correlation of Oscillation Frequencies of Fluorescence Bands and Fluorescence Transitions.

It now remains to be shown that the oscillation frequencies of the fluorescence bands of anthracene and phenanthrene may be represented as linear functions of the numbers of fluorescence transitions occurring in the systems of dynamic equilibria of their electromers. The existence of such a relationship is to be expected since the oscillation frequencies of the absorption bands of benzene chloro- and bromo-benzene and of naphthalene have been represented as linear functions of the numbers of absorption transitions occurring within the systems of equilibria of their electromers. This relationship found expression in the equation for a straight line, y = sx + b, in which y is the oscillation frequency and x the number of absorption transitions.

The oscillation frequencies of the fluorescence bands of anthucene and phenanthiene have been calculated from the values for the wave-lengths as determined by Elston and are embodied in the following table .-

		Anthracene		I'honanthrena		
		λ	ıβλ	λ	T/A	
One Two Three Four	•	390 Mt 112 Mt 135 Mt	2315 2410 2564	432 ын 415 ын 390 ын 360 ын	2315 2410 2564 2775	

The values for bands One, Two, and Three are identical for anthracene and phenanthrene.

Heretofore the supposition has been made that all of the absorption transitions within a given system were synchronous with the light waves of highest oscillation frequency. Accordingly it may now be assumed that the maximum number of fluorescence transitions, namely, twelve in the anthracene system and twenty-three in the phenanthrene system, are respectively involved in the production of the anthracene fluorescence band Three (frequency 2564) and the phenanthrene fluorescence band Four (frequency 2778) How are the numbers of fluorescence transitions corresponding to, and functioning as the origin of, the oscillation frequencies of the remaining fluorescence bands of anthracene and phenanthrene to be determined?

In the following Fig. 16, the oscillation frequencies are indicated on the Y-axis and the numbers of fluorescence transitions on the X-axis.

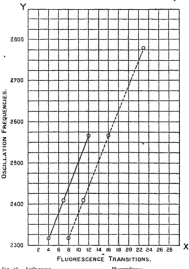


Fig. 16 -Anthracene -Phenanthrene

An examination of the tabulated groups of transitions of the anthracene system shows that four (4) is the smallest number of fluorescence transitions comprising a group. If the vibrations of four such transitions are assumed to be synchronous with the lowest oscillation frequency (2315) of the anthracene fluorescence band One, and the vibrations of the maximum number of twelve transitions be synchronous with the highest oscillation frequency (2564) of anthracene fluorescence band Three, the points (4, 2315) and (13, 2564) will determine the direction of a straight line, Now note that this straight line intersects the perpendicular from the frequency value 2410, of band Two of anthracene, at a point which has a corresponding value on the X-axis equal to the whole number (7). Hence the oscillation frequencies 2315, 2410, and 2564, of the three fluorescence bands of anthracene may be represented as a linear function of the numbers of fluorescence transitions, 4, 7, and 12, respectively. The equation for the straight line which most nearly correlates these numbers with the corresponding frequencies, as determined by the method of least squares is-

$$y = 310918.r + 31912959$$

in which y is the oscillation frequency of a given fluorescence band and x the number of fluorescence transitions functioning as its origin.

Does a similar relationship exist between the oscillation frequencies of the fluorescence bands of phenanthrene and a series of whole numbers of fluorescence transitions? Since the frequencies of the bands One, Two, Thires, are identical for both anthracene and phenanthrene, it would naturally follow that the several whole numbers corresponding to the frequencies of the three bands of phenanthrene should differ by a constant quantity from the numbers 4, 7, and 12, which correspond to the frequencies of the three fluorescence bands of anthracene. Such a relationship could only be fulfilled when the expression for the linear function takes the form of an equation for a staight line parallel to the line for authracene. In other words, the anthracene and phenanthrene lines must be parallel.

It has already been assumed that the twenty-three fluorescence transitions of the phenanthrene system correspond to the oscillation frequency 2778 of the phenanthrene fluorescence band Four, Now if a straight line be drawn through the point (23, 2778), parallel to the anthracene line (Fig. 16) it is remarkable that practically intersects the perpendiculars from the frequency values

on the Y-axis at points which have a corresponding value on the X-axis equal to the whole numbers 8, 11, and 16. In other words, the oscillation frequencies, 2315, 2410, 2564, and 2778, of the phenanthiene fluorescence bands may be represented as a linear function of the numbers of fluorescence transitions, 8, 11, 16, and 23, respectively. The equation for the straight line which correlates these numbers with the corresponding frequencies, as determined by the method of least squares.

$$y = 308255x + 20697790$$

in which y is the oscillation frequency of a given band, and x, the number of fluorescence transitions functioning as its origin.

The deviations of the calculated or theoretical values from the experimentally determined values of the oscillation frequencies of the fluore-cence bands of anthracene and phenanthrene are embodied in the following tables —

Anthracene · $y = 31'0918\pi + 2191'2959$

A B		С	D		
4	2315	2315 663	+ o 663		
7	2410	2408 938	I o61		
12	2564	2564 397	+ o 397		

Phenanthrene . y = 30'8255x + 2069 7790.

A	В	с	D		
8	2315	2316'363	+ 1'383		
II	2410	2408 859	- 1'141		
I6	2564	2562 987	- 1'013		
23	2778	2778'765	+ 0 765		

In the above tables columns (A) contain the theoretically determined numbers of fluorescence transitions represented by (x) in the linear equations for anthiacene and phenanthrene; columns (B), the experimentally determined oscillation frequencies of the fluorescence bands represented by (y); columns (C), the frequencies as calculated from the anthracene and phenanthrene equations. Note that the deviations, in columns (D), of the calculated from the actual oscillation frequencies are so small that they may be attributed to errors in experimental observation.

These results substantiate the hypothesis that the oscillation frequencies of the fluorescence bands of both anthracene and phenanthrene may be represented as linear functions of the numbers of fluorescence transitions involved in the systems of dynamic equilibria of their respective electromers. This fulfils the third object of the present chapter.

It may now be of interest to compare Elston's electronic explanation of the fluorescent spectra of anthracene and phenanthrene with the hypothesis of fluorescence transitions. Elston states that if we assume fluorescence to be produced by a system of electrons within the molecule, then, in order to account for the fluorescent spectrum of anthracene (or phenanthrene) vapour consisting of several bands superposed upon a continuous spectrum, we may consider either (1) that the electrons corresponding in period to the several bands are more numerous than those which give rise to the weaker continuous parts of the fluorescence, or (2) that the former are set in more violent vibration. It may also be assumed (3) that the system of elections is so intimately connected in its parts that, when disturbed by the exciting light in any manner, all of the electrons in the system are set in vibration. If the disturbance of the system takes place through an intermediary "luminophore" then this luminophore undoubtedly consists of a connected system of electrons whose periods correspond to those of the absorption spectrum of the vapour

The proposed hypothesis of absorption and fluorescence transitions does away with the assumption of the "luminophore" or any other group of atoms, as the cause of fluorescence, and in its place presents a definite picture of the manner in which a system of elections within the molecular structure of organic compounds may function so as to produce not only the absorption, but also or elections or of light. All of the several systems of elections or electioners are intimately connected through the centric electromer and centric rearrangements. If the system of election in a centric electromer be disturbed by the exerting light, phasotropic electioners are formed and thereby make possible the systems of dynamic equilibria which involve absorption and fluorescence transitions. The existence of certain groups of transitions, the vibrations of which are synchronous with light waves of a definite oscillation frequency, constitutes the origin of

absorption and fluorescence bands, the oscillation frequencies of which may be represented as linear functions of the numbers of absorption and fluorescence transitions respectively involved in their modulation.

G. Other Interpretations by Means of Absorption and

Other phenomena may be interpreted by the hypothesis of absorption and fluorescence transitions. Stark and Meyer 144 have observed that for numerous benzene derivatives the regions of absorption lie adjacent to or overlap the regions of fluorescence Nichols and Menut 145 also note that the broad continuous bands of fluorescent substances are always associated with a broad absorption band usually overlapping the fluorescence band on the side toward the violet, and that the absorption spectrum of the manyl salts consists of a series of hands precisely similar as regards their arrangement and number to the bands of the fluorescence spectrum. In the fluorescent spectrum of anthracene Elston observes that the bands are superposed upon a continuous region extending from 365uu to 470uu, while the absorption spectrum extends continuously from about 400µµ to some point beyond 32544. This relationship between the regions of absorption and fluorescence may readily be explained by the bypothesis of absorption and fluorescence transitions from two standpoints (1) A fluorescence transition, as previously defined. occasions absorption as well as the emission of light (2) A given type of electromer may often function both in an absorption transition, and in a fluorescence transition as is apparent in the tabulations of transitions on pages 241 and 243. Consequently the periods of vibration of these related absorption and fluorescence transitions must likewise be related. This phase of the subject ments further investigation and development

Finally, the relationship between fluorescence and phosphorscence, or luminescence, should be briefly considered. Wood 146 states that a satisfactory theory of fluore-scence must fulfil three requirements: (1) It must distinguish between inedia which fluoresce, and those which do not. It must explain (2) the change in wave-length, and (3) the increase in duration of the emission, which is the phenomenon of phosphorescence. Wood further states that at orsent there is no satisfactory theory of fluorescence. The first two requirements of a satisfactory theory have been fulfilled in the development and application of the hypothesis of absorption and fluorescence transitions. The third requirement, namely, the explanation of phosphorescence may also be interpreted in terms of fluorescence transitions on the generally accepted assumption that the energy of the absorbed light is stored in the substance in the form of potential energy of the atoms Now when the exciting source of light is removed, this potential energy, in order to be retransformed into radiant energy, causes a change in the positions of the electrons in the centric electromer. A disturbance of the condition of equilibrium in a centric electromer causes it to pass into its phasotropic electromers, and thereby a system of dynamic equilibrium is established which will involve fluorescence transitions. In other words, when the exciting source of light is withdrawn there is a continuation of the emission of light from the substance due to the persistence of fluorescence transitions which serve as the medium through which the potential energy of the absorbed light is transformed into fluorescent light.

A tentative explanation of the phosphorescence of mineral salts may also be given in terms of fluorescence transitions. It has been shown by several investigators that the action of ultraviolet light upon mineral salts causes dissociation and it has been assumed that after removal of the substance from the exciting source, phosphorescence is produced as the result of the recombination of the previously dissociated radicals. Now

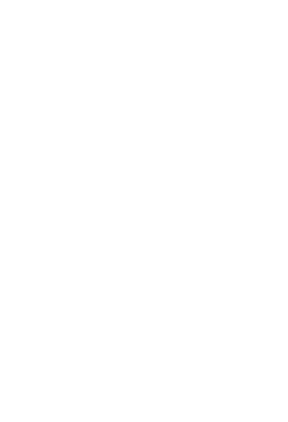
dissociation of a salt, MX, either into its ions M and X, or molecularly into M and X, cannot occur unless there be a disturbance of the relative positions of the electrons and systems of electrons which constitute the radicals M and X (see p. 153). Hence the assumption may be made that the return to the undissociated condition, MX, is accompanied by changes which concepond in some way to fluorescence transitions.

While it is possible to give a definite picture of the rearrangements of valence electrons taking place within the molecules of such compounds as benzene, naphthalene, anthracene, and phenanthrene, it is not yet possible to extend the picture to the systems of electrons within an atom. The constitution of the atom is yet quite an enigma. Therefore it is probable that the knowledge of the relationship between absorption and fluorescence phenomena and chemical constitution will be advanced chiefly through the study of the dynamic equilibria of the electromers of various organic and inorganic compounds. The absorption transition and fluorescence transition hypotheses cannot be applied to the atoms of elements because their electronic formula or structure is as yet unknown. Perhaps a study of the absorption and fluorescent spectia of the elements from the point of view of the numbers of contraples-diplex transitions which possibly function as the origin of the absorption and fluorescence bands may shed some light upon the problem of the electronic constitution of the atom.



PART IV.

METAL-AMMINES, BIBLIOGRAPHICAL REVIEW AND GENERAL CONCLUSIONS.



CHAPTER XXII.

THE CONSTITUTION OF THE METAL AMMINES

A. The Status of the Problem.

THE foremost problem of the organic chemist is generally conceded to be that of the constitution of benzene and substitution in the benzene nucleus. Another perplexing problem, of lesser importance, but of equal interest, particularly to the morganic chemist, is that of the constitution of molecular compounds, notably the metal-ammines

I W Mellor 147 states that "the attempt to distinguish molecular from atomic compounds, by structural formulæ based upon ordinary valencies deduced from the manifestations of the has not been successful". The simple atomic compounds. significance of this statement is fully realized after one has attempted to comprehend and to correlate the various theories that have been proposed to explain the constitution of the metal-ammines. Stewart 148 has presented a cutical review of the foremost theories (as applied to the cobalt-ammines) notably those of Blomstrand, Jorgensen, Weiner, Friend, Bakei, and Ramsay. Hence a comprehensive review of these theories will not be attempted. Stewart also concludes that the whole question of the constitution of the metal-ammines is at present in a very unsettled condition.

On the other hand, Sir William Ramsay 119 maintains that no theory of valency would be acceptable if it did not attempt to assign structural formulæ to the metal-ammine compounds and to correlate their properties with their formulæ. Hence it is the purpose of the present chapter to attempt to apply the electronic conception of positive and negative valence as developed in this monograph to the constitution of the chief metal-ammines. namely, the platinous-ammonia, the platinic-ammonia, and the cobaltic-ammonia compounds.

It should be noted here that the theories of Friend and 257

Ramsay, and a later scheme of Nelson and Falk 150 are based upon the electron theory Friend's system of formulation fails in its extension to the hexammine compounds since there is no valence to bind the hexatomic ring of six ammonia molecules to the remaining atoms of the compound molecule which are left suspended in space, so to speak. In Ramsay's electronic formula the ammonia molecules are singly bound to the central metal atom by a double bond of the contraplex type, i.e., the nitrogen atom of ammonia simultaneously gives to and receives from the metal atom an electron. This is the bringing into play of the so-called latent valency of Friend, identical with the neutral affinity of Spiegel and the electrical double valence of Arrhenius. Further, in Ramsav's electronic formulæ an ionizable atom or radical is attached to the nitrogen atom of an ammonia molecule which nitrogen atom is then bound by a single valence to the central or nuclear metal atom. This ionizable radical functions negatively. Nelson and Falk base then formulæ on the Weiner co-ordination types There are no new conceptions in the Nelson and Falk method of assigning electronic formulæ to the metal-ammines. The ammonia molecules and the ionizable radicals are united according to Ramsav's ideas

In the present application of the electronic conception of valence to the constituent atoms of the metal-ammines, the Werner co-ordination formulae and the various kinds of valency postulated and embodied therein, will not be considered. The chief reason for the rejection of the Werner co-ordination formulae is due to the fact that the Werner-Joigensen controversy, which extended over a period of eight years, has demonstrated that little can be gained by the wide departures of Weiner from the old and well-tried system of valency as developed in relation to organic chemistry. Therefore, in the present chapter, the electionic conception of valence will be applied only to such structural formulae for the metal-ammines as are in hairmony with the principles of valency commonly employed in the writing of the structural formulae of carbon compounds,

The structural and electronic formulæ herewith proposed will be derived through (1) the development of a few simple rules relative to the distribution of the positive and negative valences, and (2) the application of these rules to the empirical formulæ of the typical metal-ammines. To this end, the rational names and the empirical formulae of the three foremost groups of metal-ammines will be tabulated as follows: (A) the platinous-, (B) the cobaltic-, and (C) the platinic-ammonia compounds

TABLE A.

PLATINOUS-AMMONIA COMPOUNDS.

Rational Name	Ismpirecal Formula
1 Tetrammine-platinous salts	. [Pt(NH,)]X. [Pt(NH,)]X
2, Triammine-plaunous salts .	[Pt(NH ₂) ₁ X]X
 Diammine-platinous compounds. 	[Pt(NH ₂),X] (two isomer

TABLE B.

Conaltic-Ammonia Compounds

	Hexammine-cobaltic salts .		[Co(NII.) _a]X.
	Pentammme-cobaltic salts .		[Co(NIL) ₅ X]V,
	Tetrammine-cobaltic salts .		[Co(NH ₃),X.7\(\) (two isomers)
1.	Triainmine-cobaltic compound		[Co(NH) N]

TABLE C

PLATERC-AMMONIA COMPAUNDS.

	Hexammine-platinic salts ,		[Pt(NH _d) _d]X _d [Pt(NH _d) _e X]X _e (unknown)
	Pentamm ne-platime salts		(Pt(NH ₃), X ₁ X _* (unknown)
3.	Tetrammine-platinic salts		(P((NH), X))X
4	Triammine platinic salts		[Pt(NH), Y, IX [Pt(NH-), X, IX
5	Drammine-platinic compounds		[Pt(NH _a),N _a] (two isomers)

B. Fundamental Principles Pertinent to the Electronic Formulæ of the Metal-Ammines.

A survey of the three groups (A, B, and C) of metal-ammonia compounds reveals two remarkable relationships which constitute the basis for definite rules to be employed later in writing the electronic formulæ of metal-ammines. Relative to the development of the first relationship, note that in Table A there are three classes of platinous-ammonia compounds. In platinous compounds, platinum is bivalent In Table B there are four classes of cobaltic-ammonia compounds. In cobaltic compounds. cobalt is tervalent Lastly, in Table C, there are five classes of platinic-ammonia compounds In platinic compounds, platinum is quadrivalent. In other words, when the valence of the metal atom of the metal-ammonia compound is (n), then there are (n + 1) classes of metal-ammines. In this connection recall the electronic valence rule previously developed and illustrated, namely, that when the valence of an atom is (n), that atom may function in electronic formulæ according to (n + 1)electionic types. The existence of this remarkable relationship between the numbers of classes of metal-ammines and the electronic valence rule leads to the immediate conclusion that the number of classes of the metal-ammines of a given metal is related directly to the number of the electronic types in which said metal atom may function. More specifically, the three classes of platinous-ammonia compounds noted in Table A correspond to the three electronic types :-

The four classes of cobaltic-ammonia compounds of Table B correspond to the four electronic types:—

Lastly, the five classes of platinic-ammonia compounds of Table C correspond to the five electronic types:—

The second relationships based upon the fact that some of the metal-ammines display ionogen properties while others do not. A definite correlation between the electronic type of the metal atom and the number of ionizable atoms or radicals (X) which exist in the compound is found in the fact that the number of positive valences of the metal atom is equal to the number of the megative ronizable groups (X) This principle makes it possible to relate a particular electronic type of the metal atom to a cone-sponding type of a metal-aminine. For example, in the platinous-aminine group, the tetrammine and the triammine salts display respectively two and one dissociable negative radicals (X). These numbers are equal to the numbers of positive valences of the metal atom, namely, two and one, which correspond respectively,

to the electronic types $Pt \stackrel{\cdot}{+}$ and $Pt \stackrel{\cdot}{-}$. In this connection the diammine platinous compound displays no longen properties and, accordingly, its platinum atom, having no positive valences, conforms to the electronic type $Pt \stackrel{\cdot}{-}$.

In the same manner, and according to the same principle, the four groups of cobaltic-ammines which display three, two, one, and no ionizable negative radicals, embody respectively cobalt atoms of the electronic types,

In other words, the number of negative nonizable atoms or radicals of the metal-annune is equal to the number of positive values of the metal atom of the metal-annune. This relationship, found to hold for the platmous-, cobaltic-, and platinic-annune compounds, constitutes the second principle involved in the writing of the electronic formulae of the inetal-annunes.

Before postulating the rules for writing the electronic formulaof metal-ammines some account must be given of the manner in which the ammonia molecules are to be distributed. It will be remembered that Blomstrand, and later Jorgensen, assumed a division of the ammonia molecules of the metal-ammines into chains by virtue of the ability of the nitrogen atom of ammonia to pass from its tervalent state to the quinquevalent condition. The pentamine-cobalité salt was written thus.—

The non-ionizable chlorine atom is directly united to the cobalt atom; the two ionizable chlorine atoms are united to nitiogen and thus have the same ionogen properties as chlorine in aimmonium chloride. Note that the above formula, comprising five aimmonia molecules, has one chain of four aimmonia molecules. Blomstrand and Joigensen have advanced no reasons indicating how many aimmonia molecules should be embodied in a nitrogen chain. Hence the above pentainnine salt could be represented just as well by the following isomeric formula.

which contains chains of two and three ammonia molecules. In other words, the Blomstrand-Jorgensen method of assigning structural formulæ to the metal-ammines permits of more isomers than the facts warrant. To overcome this difficulty, which is also likely to be encountered in writing the electronic for mulæ of metal-ammines, there must be some limitation and definition of the number of ammonia molecules to be embodied in a nitrogen chain.

To this end, a guiding principle is found in the fact that some metal-ammonia compounds, although generally stable, indicate a proneunced tendency to love ammonia molecules in pairs. For example, in an atmosphere of ammonia at 760 mm, piessure, the compound Zn(NH₂),Cl₂ decomposes at 59°, losing two molecules of ammonia, and in the same way the resulting compound Zn(NH₂),Cl₂ also loses two molecules of ammonia at 89°5' yielding Zn(NH₂),Cl₂. Finally, this compound decomposes at 26°9' with the loss of its remaining two molecules of ammonia. These transitions are summaized thus,—

$$Zn(NH_2)_iCl_2 \rightarrow Zn(NH_2)_iCl_2 \rightarrow Zn(NH_3)_iCl_2 \rightarrow ZnCl_2$$

Further evidence for the apparent association of groups of two molecules of ammonia in the metal-ammines is found in the fact that Magnus' Green Salt, [(NH₂)Pt]PtCl₃], is made by the action of ammonia upon platinous chloride, PtCl₂. When boiled with ammonia it yields tetrammine-platinous chloride, [Pt(N H₂)]Cl₃. The tetrammine salts when heated also lose two molecules of ammonia yielding the symmetrical diammine compounds of general formula [Pt(NH₂)]Cl₃]

The pronounced tendency, noted in the foregoing facts, for ammonia molecules to enter into combination or to be eliminated from combination m pairs, warrants the assumption that two constitutes the number of ammonia molecules naturally occurring in a chain of ammonia molecules. Therefore, in postulating rules for writing the structual and electronic formule of the metal-ammines, it will be assumed that not more than two molecules of ammonia will be embodied in a single nitrogen chain of chain of ammonia molecules.

C. Rules for writing Structural and Electronic Formulæ of

The rules for writing the structural and electronic formulae of the metal-ammonia compounds noted in Tables A, B, and C (p, 259) are as follows —

 Write all of the possible electronic types of the metal atom of the metal-amminos of a given metal. When the valence of the metal is (n) there are (n + 1) electronic types and (n + 1) classes of metal-ammines.

(2) Since the number of negative ionizable atoms or radicals (X) of the metal-ammine is equal to the number of posture valences of the metal, attach to each positive valence of the metal atom (M) a chain consisting of two molecules of ammonia and one atom or radical (X) which is ionizable and functions negatively, thus —

This electronic formula for the chain scheme may be abbreviated by indicating the polarity, omitting the valence line, and letting (a) represent a molecule of ammonia, thus:—

In applying the above rule (2) it is quite evident that no more molecules of ammonia may be written into the electronic formula than actually exist in the empirical formula. However, as far as the number of ammonia molecules will permit, this rule should be applied to each positive valence of the metal atom.

If the metal-ammune contains an add number of ammonia molecules it will be impossible to assign all of them in pairs. Accordingly, the remaining odd number of ammonia molecules may be united singly to any negative valences of the metal atom, and these single ammonia molecules will in turn be united to an atom or radical X which functions positively, thus:—

It should be noted that when X is an acid radical, for example, chlorine or bromine, and is positive, it manifests little, if any, tendency to ionize. This is partly analogous to the situa-

tion presented by acids of the type $H \stackrel{*}{\longrightarrow} X$ and $H \stackrel{*}{\longrightarrow} O \stackrel{*}{\longrightarrow} X$. The tendency for X when negative to function as an anion is very pronounced; but when X is a non-metal and positive, its tendency to function as a cation is slight. (See Chapter VI on ionic amphoterism.) On the other hand, if X is a metal it tendency to function positively as a cation is quite masked Apropos of the preceding statements the third rule is as follows:

(3) To each negative valence of the metal atom attach a chain consisting of one molecule of airmonia and one atom or radical X which will function positively, thus:—

$$M - + a - + X$$
.

If, in the application of rules (2) and (3) all of the available molecules of ammonia are distributed before all of the positive or negative valences of the metal atom are disposed of, the latter are united directly (without the interposition of ammonia molecules) to any remaining atoms or radicals, X. If the remaining valence of the metal is positive, then X is negative, thus M + - X° If the remaining valence of the metal is negative, then X is positive, thus M - + X. This procedure maintrains the principle previously noted that the number of the positive valences of the metal atom equals the number of negative atoms or radicals, while the number of negative valences of the metal atom equals the number of positive valences of the metal atom equals the number of positive valences of the metal atom equals the number of positive valences of the metal atom equals the number of positive valors or radicals.

Before applying the above rules, the existence of isomers of the metal-ammines should be noted. Then existence is correlated with the following rule (4):—

(4) Isomers are possible and are accounted for whenever in the application of the preceding rules there remain for disposition one pair of negative valences of the metal atom, two molecules of annionia, and two atoms or radicals which function positively. These may be disposed according to two structural electronic formulae each of which (1) and (II), persents an isome.

Note that isomer I takes care of the negative valences of the metal atom according to rule (3). The possibility of another formula, isomer II, is somewhat exceptional in that a chain of two molecules of ammonia is united to a negative valence of the metal.

Isomers of the following metal-ammines are known .--

In these compounds the electronic types of the metal atoms are respectively as follows:—

In the application of the rules for writing the electronic formule of these compounds, there will remain for disposition in each, one pair of negative valences of the metal atom, two molecules of ammonia and two atoms, $X_{\rm p}$ which are positive Hence the co-ultions exist for the two isomeric formulae in conformity with rule (4) and the empirical facts

D. Applications of the Rules. *

The preceding rules (i-4) described and qualified, are based upon electronic principles established in conformity with empirical data. They are, therefore, not to be regarded as arbitrary or hypothetical, but rather as affording a method of writing both the structural and the electronic fointile of the metal-ammines. These rules will now be applied in the writing of the electronic formulae of the platinous, the cobaltic, and the platinic-ammonia compounds. Their fointile are recorded in tabular foim. First, note that the empirical formulae (taken in consecutive order from the preceding Tables A, B, and C of the metal-ammonia compounds) are immediately followed by the electronic formula or type of the metal atom in the given metal-ammine. These are followed by their electronic formula which are derived by an application of the foregoing rules to the positive and negative valences of the metal atom.

(1) [Pta, Va]X

(5) [Pta₂X₄] (two isomers, I and II,)

THE ELECTRONIC CONCEPTION OF VALENCE TABLE A . PLATINOUS-AMMONIA COMPOUNDS. (t) | Pta,]X., 1 - 4 1 - 4 + - 1 Pt 4 ^ (2) [Pta₁X]X (3) [Pta,X.] (two isomers, I and II) TABLE B. COBALTIC-AMMONIA COMPOUNDS + - a + - a ! - 1 .. (1) (Coa.3X) Co + - a + - a + - 1 - a + - a + - 1 (2) [Coa5X]X2 Co + - a + - a + - x Co - + x (3) [Coa, V.]X (two isomers, I an l II) Co - + a - + x (1) [Cua, X,1 -+4-+1 TABLE C PLATINIC-AMMONIA COMPOUNDS + - a + - a + - x + - a + - a + - + (t) [Pta₆]X₄ +- a + - a + - 2 (Formula cannot be written (2) [Pta, X]X, in conformity with rules) (unknown) Pt + - a + - a + - 1 (3) [Pta₁X₂]X₂ - + 1

+ - a + - a + - 1 - + a - + x

- + r - + r - + a - + 1

II Pt

.

It is particularly noteworthy that no pentammine platinic salt of formula (2), Table C. namely [Pta, X]X2, is known. The rules for writing the electronic formulæ are not applicable to a metal atom of the type Pt + because five molecules of ammonia

cannot be attached in groups of two to each of the three positive valences of the metal An electronic formula may be written,

but it is an exception to the rules which are applicable to each of the known metal-ammines. The fact that this pentammine platinic salt does not exist, and that an electronic formula in conformity with the rules cannot be written, lends support to the proposed system of writing the structural and electronic formulæ of the metal-ammines.

It is also significant that the metal-ammine salts undergo electrolytic dissociation, that is, they ionize in aqueous solution, in conformity with the electronic formula. When X is negative. it functions as an anion. The remaining part of the molecule constitutes the cation, or positively charged complex radical For example, the ionic dissociation of the pentammine cobaltic salt is represented thus :-

These electronic formulæ suggest the possibility of explaining another type of electrolytic dissociation in which X is positive and functions as the cation. The remaining part of the compound is the complex negative radical or anion (M = neutral atom):

E. Metal-Ammines and Complex Salts: Transition Series.

This leads to an interpretation in terms of electronic formulæ of the series of seven cobaltic compounds in which the valence of the complex radical changes progressively from three positive to three negative. The empirical formulæ of these compounds,

recorded in the order of the noted transitions in valence of their complex radicals, are as follows:—

 $[Co(NH_4)_d]$ Cl_2 ; $[Co(NH_4)_s(NO_s)]$ Cl_2 , $[Co(NH_4)_s(NO_s)_2]$ Cl, $[Co(NH_4)_s(NO_s)_d]$, $K_2[Co(NH_4)_s(NO_2)_s]$; $K_3[Co(NO_s)_a]$.

Before tabulating the electronic formulæ of this series of cobaltic compounds, the question of the disposition of the NO $_2$ groups must be considered. It should be recalled that the tadical NO $_4$ may function either negatively or positively, according as it is

looked upon as a derivative of H. NO, or HO. NO, respectively. Furthermore, NO, as a compound, or as an independent molecule. has been shown (p 153) to function as a "free radical" in which the valence of its nitrogen atom is four. Accordingly, through the gain or through the loss of an election it becomes a negative or a positive radical respectively. Since it may function as an independent molecule, there is nothing to preclude the assumption that chains of molecules of NO. may exist and function in the same manner as do chains of molecules of ammonia in the metal-ammines. Such chain formation is attributed to the canacity of the nitrogen atom, either of NH, or of NO, to part with and acquire simultaneously an electron, i.e., to develop a free positive and a free negative valence. Hence the following partial schemes show how (a) a negative atom may be at the end of a chain comprising two molecules of NH,, and (b) how a positive atom may be at the end of a chain containing two molecules of NO:--

(a)
$$M + - NH_3 + - NH_3 + - CI$$
,
(b) $M - + NO_3 - + NO_3 - + K$,

These arrangements maintain the rule that for every negative valence of the metal there shall be a positive atom, adical, or on; and, for every positive valence of the metal there shall be a negative atom, radical, or ion.

The first column of the following table presents the empirical formulæ of the series of codalitic compounds in question. The second column notes the magnitude and the polarity of the valence of the complex radical. Column three indicates the electronic type of the metal, followed, in the last column, by the electronic formula which is derived through an application of the previously developed rules and principles.

Number of Com- pound	Empirical Formula	Valence of the Complex Radic il	Electrome Type of Metal	Ficetronic I ormula
(1)	[Co(N H ₀),]CI,	3 +	Co + +	+ - NH ₁ + - NH ₁ + - CI Co+ - NH ₁ + - NH ₁ + - CI F - NH ₁ + - NH ₃ + - CI
(4)	[Co(NH4)4(NO [®])]Cl [®]	2 +	Co +	+ - NH ₁ + - NH ₂ + - CI Co+ - NH ₁ + - NH ₄ + - CI - + NH ₁ - + NO ₂
(3)	[Co(NH ³) (NO ³)]CI	x 1	Co -	+ - NH ₁ + - NII ₁ + - Cl Co - + NII ₁ - + NO, - + NII ₁ - + NO,
(1)	[Co(NII),(NO),]	D	Co -	-+NII,-+NO, Co-+NII,-+NO, -+NII,-+NO,
(5)	[Co(NO ₂) ₀]K,	3 -	Co -	-+NO ₃ -+NO ₂ -+K Co-+NO ₃ -+NO ₃ -+K -+NO ₃ -+NO ₂ -+K
(6)	[Co(NO ₂),(NII ₃)]K ₂	3	Co -	-+NO ₂ -+NO ₂ -+K Co-+NO ₂ -+NO ₂ -+K +-NH ₁ +-NO ₂
(7)	[Co(NO ₂),(NH ₄) ₂]K	1 -	Co +	-+NO ₂ -+NO ₂ -+K Co+-NH ₁ +-NO ₂ +-NH ₁ +-NO ₂
(4)	[Co(NO ₃) ₁ (NH ₄) ₃]	0	Co +	+ - NH ₁ + - NO ₂ Co+ - NH ₁ + - NO ₂ + - NH ₂ + - NO ₃

A critical survey of the above table shows that the negative chlorine atoms (or ions) of compounds (1), (2), and (3), and the positive potassium atoms (or ions) of compounds (5), (6), and (7), are duly indicated in the electronic formulæ of the respective This correlates the ionogen properties of the compounds. compounds with their electronic formulæ.

The electronic formulæ of compounds (1), (2), (3), and (4) are in complete accord with the postulated rules and accordingly are perfectly similar, both structurally and electronically, to the formulæ of the cobaltic-ammonia compounds (1), (2), (3), and (4), respectively, of Table B (p 266) Compounds (5), (6), and (7), however, embody NO, molecules, and NO, radicals, as previously explained. It should be noted that compounds (1) and (5), (2) and (6), (3) and (7), and (4) and (8), are perfectly similar structurally; but electronically there is a complete reversal of the polarity of the valences not only of the cobalt atoms, but also of the remaining molecules and radicals which compuse the compounds.

In this connection, it is remarkable that structurally identical, but electronically opposite formulæ, (4) and (8), are electronic isomers of the compound [Co(NH_a),(NO₃), which is a nonelectrolyte. In other words, formulæ (4) and (8) are electromers of the cohaltic-trinitro-triammine. The existence of electromers of organic compounds has been demonstrated, i.e., the reactions of certain compounds can be explained only by assuming the existence of its electromers, each of which enters into a definite chemical reaction vielding its own specific derivative. The existence of electromers of inorganic compounds such as formulæ (4) and (8) is probable. Such electromers may exist in tautomeric equilibrium (clectronic tautomerism) or, if the properties of the compound so indicate, only in one electronic form. Of the two possible formula, (4) is the more likely because its NO. radicals are positive and it is a non-electrolyte. This accords with the well-known fact that when the radical NO. is positive. it does not tend to function as an ion.

The principles and rules presented in this chapter for writing the structural and electronic formulæ of the metal-ammines might be extended almost indefinitely to various other series of complex inorganic or molecular compounds, but enough has been given to suggest the possibility and the method of extension. Whether the types of formulæ here proposed are more consistent and more significant than previously proposed formulæ for the metal-ammines is, of course, a question. One claim, however, is made, namely, that the proposed electronic formulæ have not departed from the thoroughly established principles which underlie the writing of the structural and graphic formulæ for each or compounds. Furthermore, the electronic formulæ definitely qualify the valences as positive or negative and thereby effect more complete correlation with chemical properties.

Since the electronic conception of positive and negative valence as applied to the constitution of benzene and many of its derivatives has made it possible to explain and to correlate many hitherto inexplicable and unrelated chemical and physicochemical phenomena, it is to be hoped that the extension of the

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electronic conception of walence, as herewith applied to the constitution of the metal-ammines, may lead to a more complete development of the structural and electronic formula: of both molecular and atomic compounds and thus bring them into a unified system.

CHAPTER XXIII.

BIBLIOGRAPHICAL REVIEW.

A REVIEW of the contents of the many articles relating either directly or indirectly to the electronic conception of valence would require another volume. If, however, a review is limited to the published applications of the electronic conception of valence in so far as they relate to the interpretation and conselation of chemical and physico-chemical phenomena, it may be embodied conveniently in a single chapter. Accordingly the present chapter is limited to a bibliographical and chromological review of the applications of the electronic conception of valence presenting. (1) the name (or names) of the author of the given article, (2) the title of the article, (3) the reference to the journal in which the article appeared, and (4) a brief abstract of the contents of the article.

A. Bibliographical Review of Published Applications.

The first applications of the electronic conception of valence to the interpretation of specific chemical phenomena by means of electronic formula were presented by H S. Fiy in a paper isad before the Cincinnati Section of the American Chemical Society (January 15, 1908) entitled "An Hypothesis relative to the Constitution of the Benzere Nucleus an Application of the Corpuscular Atomic (Electronic) Conception of Positive and Negative Valences to the Constituent Atoms of Benzere," J. Amer. Chem. Soc. 30, 34 (1908). In this paper, Fry elaborated the electronic conception of positive and negative valence (originally suggested by Sir J. J. Thomson's work, Electratia and Matter), and proposed the new conceptions and terms electronic, electronic tautomasium, and the electronic formula of benzene with an expolanation of the Biowa and Glusson rule.

In the following year J. M. Nelson and K. G. Falk, also

basing their views upon Thomson's hypothesis that the linkages between atoms in a compound are caused by the transfer of corpuscles, applied same to a number of facts chiefly from organic chemistry "The Electronic Conception of Valency in Organic Chemistry," School of Mines Quarterly, 30, 179-198 (1909) See also I. Amer, Chem Soc., 32, 1637-1654 (1910), which practically embodies then first paper and gives a general discussion of aliphatic carbon and nitrogen compounds containing single, double, and triple bonds; compounds containing double bonds between unlike atoms; partial valence, and complex inorganic salts In their summary, Nelson and Falk state that "all cases of isomerism connected with the presence of a double bond, whether between like or unlike atoms, have been referred to the direction of the valences of the double bond, instead of to spatial configurations as heretofore. The existence of certain isomers and the explanation of some hitherto unexplained reactions have also been referred to the direction of valences The existence of 'partial valence' is shown to follow from the electric charges in a molecule"

H. S. Fiy. "Die Konstitution des Benzols vom Standpunkte des kopuskular-atomistischen Begriffs der positiven und negativen Weitigkeit. I Eine Interpretation der Regel von Crum Biown und Gibson" Zeitsch. physikal. Chem., 76, 385-397 (1911). The principles presented in this papea are embodied and fürther developed in Chapters II., VII., IX., and X. of this monograph.

H. S. Fry: Idem. II. "Dynamische Formeln und das Ultraviolettabsorptionsspektrum des Benzols". Zeitschr physikal Chem., 76, 398-412 (1911). See Chapter XVIII.

H S. Fry Idem. III "Dynamische Formeln und das Ultraviolettabsorptionsspektrum des Naphtalins". Zeitschi physikal. Chem., 76, 591-600 (1911) See Chapter XX.

K. G Falk "Electron Conception of Valence. II The Organic Acids" J. Amer. Chem. Soc. 33, 1140-1152 (1911). Falk proposes a classification of the organic acids according to the direction of the valences by which the alpha carbon atom is combined with the other atoms of the molecule. The ionization constants (K × 10°) depend primarily upon the additive effects of the directive valences of this a-carbon atom: I. \$\frac{1}{2}C.CO_2H, ionization constants less than 001; II. \$\frac{1}{2}C.CO_2H, ionization

constants on - o.4, III. \(\xi \) C.CO.H, ionization constants greater than 2

H. S. Fiy. "A Critical Survey of Some Recent Applications of the Election Conception of Valence". J. Amer. Cleim Sur34(664-673 (1912). Fiy maintains that Falk's classification of the organic acids is incomplete because it fails to take into account the direction of the valence which binds the a-caubon atom to the carboxyl group. When this valence is considered, the theory demands eight classes of organic acids. Falk postulates only four and considers experimental data for only three classes. Furthermore the direction of the valence which binds the a-carbon atom to the carboxyl group, and which Falk ignores, is of prime import tance because it determines whether the carboxyl group functions positively or negatively, i.e., whether it corresponds electronically and chemically to the carboxyl radical in carbonic acid, or to the carboxyl radical in formic acid, or

W A Noyes. "A Possible Explanation of Some Phenomena of Ionization by the Election Theory." J Amer Chem. Sor., 34, 663 (1912). A short note also suggesting that ionization phenomena are related to the electronic state of certain atoms in the molecules of election, but the molecules of elections.

J M Nelson and K. G Falk "The Electron Conception of Valence. III. Oxygen Compounds." Communication, 8th International Congress of Applied Chemistry, 6, 212-221 [1912]. An extension of their ideas to the oxygen atoms of certain compounds.

H. S. Fiy "Einige Anwendungen des Elektronbegriffs der positiven und negativen Wertigkeit. IV Fluoreszenz: Anthiazen und Phenanthren" Zeitschr. physikal Chem., 80, 29-49 (1912). See Chapter XXI

H. S Fry Idem V. "Absorptionsspektra und dynamische Formeln von Chlor-, Brom- und Iod-benzol". Zeitsche physikal. Chem., 82, 665-687 (1913) See Chapter XIX.

W. A. Noyes. "An Attempt to Prepare Nitro-nitrogen Trichloride, an Electromer of Ammono-nitrogen Trichloride". 1 Amar. Chem Sor., 35, 767-775 (1913). Ordinary nitrogen trichloride when tutrated against assenious acid is equivalent to six atoms of chlorine per molecule indicating that the formula is Noves maintains that there should be an electromer of the formula,

nitro-nuttogen trichloride, which would hydrolyze like other nonmetallic chlorides to give nitrous and hydrochloric acids. To obtain this electromer, nitrosyl chloride and phosphorus pentachloride were brought together at 1000°-1100° to effect the reaction —

Much nitrosyl chloude temained unchanged. Chlotine and silicon tetrachloride were formed Of thirty gas mixtures analysed, twelve obtained in six different experiments gave evidence of the formation of minute traces of the electrome intronitrogen trichloride: 0 12-0 22 millimoles in a volume of about 70 cubic centimetres.

J. M. Nelson, H. T. Beans, and J. K. Falk. 1V. "Classification of Chemical Reactions.". J. Amer. Chem. Soc., 35, 1842. 1821 (1913). The authors review and define more rigorously the terms and assumptions presented in their preceding papers. As more general than oxidation, the term adduction is proposed. On the basis of the electronic changes involved, chemical reactions are classified into oxidation-reduction changes, "onium" compound formation, and simple replacement or Lexitagement. These classifications are illustrated by various electronic formule.

Two papers of ielated interest on "Valence and Tauto-mensm" should be noted One was published by W C. Bray and G. E. K. Branch, James Chem Soc., 35, 1440-1447 (1913); the other by G. N. Lewis, ibid., 35, 1448-1455 (1913) Both papers deal chiefly with polemical and hypothetical distinctions, from the electronic point of view between polar and non-polar valences

K G. Falk and J. M. Nelson V. "Polar and Non-polar Valence". J. Amer Chem Soc., 36, 209-214 (1914). The authors oppose the view of Bray and Branch that valence is sometimes polar, sometimes non-polar, pointing out that in many cases, as in the Gingand reaction, we would, if that view is correct, have the two kinds of valence appearing in the same icaction. To make such a distinction is difficult, confusing, and unnecessary.

L. W. Jones: "Applications of the Electronic Conception of Valence, Part I, Reactions among Certain Compounds containing Nitrogen. Part II The Beckmann Rearrangement." Amer. Chem 1., 50, 414-443 (1013) Iones extends the electranic valence rule of Fiv. namely, that when the valence of an atom equals (n) that atom may function in electronic formulæ in (n + 1) ways, to tervalent and to quinquevalent nitrogen atoms which present respectively, four and six electronic types the point of view of electronic oxidation-reduction reactions, many of which are intramolecular, the electronic types are displayed by and correlated with the chemical properties of amines. aldimes, nitriles, nitrile oxides, and many other nitrogen compounds. Hydrolysis is the chief means of determining the state of oxidation of the autrogen and carbon atoms. Jones maintains "that a carbon atom when linked directly to a nitrogen atom does not readily take from it negative electrons, or, in other words, is not readily reduced by it". An inspection of all of the reactions classed as examples of the Beckmann rearrangement (including the Hofmann and Curtius reactions) shows that "in every case the tearrangement is accompanied by a process of intramolecular oxidation and reduction It seems very probable that this tendency of the system of linking carbon-nitrogen to pass to one in which the carbon atom is as fully oxidized as possible. and the nitrogen atom as fully reduced as possible may be the real determining factor in the Beckmann rearrangement, and that the formation of univalent nitrogen, proposed by Stieghtz as the immediate cause, may be a mere incident, necessary, to be sure. to pave the way for this change." An appreciation of the nature and extent of Jones' applications of the electronic conception of valence to the compounds of nitiogen requires a detailed study of the published paper.

H. S. Fty: "Interpretations of Some Steroochemical Problems in terms of the Electronic Conception of Positive and Negative Valences. I. Anomalous Behaviour of Certain Deivatives of Benzene." J. Amer. Chem. Soc., 36, 248-262 (1914). See Chapter VIII.

H. S. Fry: "Positive and Negative Hydrogen, the Electronic Formula of Benzene and the Nascent State". J. Amer. Chem. Soc., 36, 262-272 (1914) See Chapters IV. and V.

Julius Stieglitz and P. N. Leach "The Molecular Re-

arrangement of Triarvlmethyl-Hydroxyl-Amines and the 'Beckmann' Reamangement of Ketoximes". J. Amer Chem. Soc., 36, 272-301 (1914). It is possible that the chloroimides obtained by Stieghtz and Peterson are electromers of the chloroimides supposed to be the intermediate products in the rearrangement of the ketoximes by phosphorus pentachloride Many rearrangement reactions are discussed in terms of the electronic valences of the nitrogen atom in relation to the positive and negative character of certain atoms and radicals. Stieglitz, in presenting a new interpretation of the rearrangement of ketoxinies, maintains that "with the change of electronic forces, the positive radical, nearest to the field of force, is lost by the now positive carbon and carried to the now negative nitrogen. Such a series of actions would account for the nature and action of the reagents used to accomplish the rearrangement (acid dehydrating agents) and it gives a rational picture of the electrical forces in play in the rearrangement of the valences of the molecule. Such a course would also account for the influence of stereoisomerism on the reairangement, if such an influence should be established as beyond doubt-the radical nearest to the electrical fields of force produced by the migration of electrons from carbon to nitrogen passing under the influence of this force to the nitrogen." Again, it should be stated that an intimate study of the published paper necessarily precedes an appreciation of the proposed interpretations.

L. W. Jones: "Electromers and Stereomers with Positive and Negative Hydroxyl". J. Amer Chem. Soc., 26, 1268-1290 (1914). Jones cites experimental facts which he believes are sufficient not only to establish a tautomeric relationship between hydroxylamine and its derivatives in the ordinary structural sense but also to confirm the belief that these compounds in many of their reactions behave tautomerically in the electronic sense, i.e., afford examples of electronic tautomerism. The paper deals in particular with: (1) structural tautomerism of hydroxylamine derivatives, (2) oxidation and eduction of hydroxylamine and its derivatives; (3) the action of hypochlorous acid and bromine upon tetlary amines; (4) an electronic explanation of these actions, (5) electromer tautomerism of hydroxylamine and its derivatives; and (6) electromers and stereomers with positive and negative hydroxyl, or

allooxyl. In the last section, Jones bloom continuely that certain decreatives of hydrocylamure, prepared by Meisenheimer, are electromers. The following quotations taken freely from a subsequent paper by Jones (Savier, Vol. 46, No. 195, 493-502) presenting this same phase of the subject, are of moment.

"In an article concenning the 'Non-Equivalence of the Five Valences of Nitrogen,' Meisenheimer describes the preparation of two isometic compounds of the type, (R),N(OCH,NOH). The first isomer was obtained by the action of (1) methyl iodide upon trimethylamine oxide and (2) subsequent replacement of iodine by hydroxyl. Thus—

(1)
$$(CHJ_3N \leq O + CHJ_1 \rightarrow CHJ_1N \stackrel{OCH_3}{\subset}$$

(2) $(CHJ_3N \stackrel{OCH_3}{\subset} + NaOH \rightarrow (CHJ_4N \stackrel{OCH_3}{\subset} + NaIH \rightarrow CHJ_1N \stackrel{OCH_3}{\subset}$

"The second isomer was secured by the action of sodium methylate upon the salt obtained by treating trimethylarylamine oxide with hydrogen chloride —

(1)
$$(CH)_1N = 0$$
 + $IICI \rightarrow (CH)_1N \nearrow OH$
(2) $(CH)_1N \nearrow OH$ + $N_AOCH_1 \rightarrow (CH)_2N \nearrow OCH_1$ + N_ACI

The two forms, (A) and (B), are identical except for the order in which the hydraxyl groups and the methaxyl groups are introduced. In (B), as Messenheimes and, the methoxyl group is linked to the 'ffith' valence,' or the one which usually engages the acid radical; while it is linked to the 'fourth' valence' in formula (A). But these two substances (structural isomers) are fundamentally different. When an aqueous solution of (A) was heated, it decomposed quantitatively according to the equation.

$$(\Lambda) (CH_2)_1 N \xrightarrow{OH} (\S) \rightarrow (CH_2)_1 N + CH_2O + U_2O.$$

The trimethylhydroxyammonium methylate (B) showed a totally different behaviour —

(B)
$$(CH_1)_{\tau}N < OH_1 (\tau)$$
 \Rightarrow $(CH_1)_{\tau}N = O + CH_2OH$,

"In addition to these compounds, Meisenheimer prepared a number of isomeric mixed dialkyl compounds with methyl, ethyl, and propyl radicals, e.g.—

$$(CH_a)_i N_i \underbrace{\begin{array}{c} OCH_i \\ \cdot OC_2H_a \end{array}}_{And} \quad (CH_i)_a N_i \underbrace{\begin{array}{c} OC_2H_a \\ \cdot OCH_a \end{array}}_{And}$$

In every case, water decomposed compounds of this type to give a textiary amine, an alcohol, and an aldehyde, but, invariably, the radical eliminated as aldehyde was the radical which occupied 'position four (4),' and the group eliminated as alcohol always occupied 'position five (5)' Meisenheime stated that he never obtained recognizable traces of the aldehyde which should have issuited if the group attached in position five had separated in that form.

"The electronic conception of valence fully explains these disputed relations by assuming that the one hydroxyl (or alkoxyl) group (4) is positive while the other (5) is negative. Thus —

$$(\Lambda) \ (CH_d)_1 N \stackrel{+}{-} \stackrel{+}{+} \stackrel{-}{-} \stackrel{+}{-} \stackrel{+}{+} \stackrel{-}{-} \stackrel{+}{+} \stackrel{-}{-} \stackrel{+}{+} \stackrel{-}{-} \stackrel{+}{-} \stackrel{-}{-} \stackrel{-}{-} \stackrel{+}{-} \stackrel{-}{-} \stackrel{+}{-} \stackrel{-}{-} \stackrel{-}{-} \stackrel{-}{-} \stackrel{+}{-} \stackrel{-}{-} \stackrel{-}{-} \stackrel{-}{-} \stackrel{+}{-} \stackrel{-}{-} \stackrel{-$$

(B)
$$(CH_1)_2N = + 0 = + H$$
 (1) \Rightarrow $(CH_1)_1N = + 0 + (CH_1 + - 0 = + H).$

"It is significant that the two oxygen atoms upon which the existence of the electromers depends are not linked directly to each other but through an intermediate atom, nitrogen; thus —

Jones maintains that this is undoubtedly responsible for the relative stability of these electromers as compared with others in which the atoms of different polarity are directly connected, e.g.—

H. S. Fry, "Interpretations of Some Stereochemical Problems in Terms of the Electronic Conception of Positive and Negative Valences II. Halogen Substitution in the Benzenie Nucleus and in the Side Chain" J. Amer. Chem. Soc., 36, 1035-1047 (1914). See Chapter XV.

J. M. Nelson and K. G. Falk. "The Electron Conception of Valence. VI. Inorganic Compounds" J. Amer. Chem. Soc., 37, 274-286 (1915) Applications are extended to Werner's conceptions. The authors, adopt the suggestions of Ramsay

relative to the electronic constitution of the cobaltic-ammines and extend same to platinic-ammines. Werner's co-ordination formula: are the structural basis for the electronic formulae of the inetal-ammines as conceived by Rainsay and later by Nelson and Palk

H. S. Fiy, "Interpretations of Some Stereochemical Problems in Terms of the Electronic Conception of Positive and Negative Valences. III. A Continuation of the Interpretation of the Brown and Gibson Rule" J. Amer. Chem. Soc., 37, 855-863 (1615). See Chapter IN.

H. S. Fry: Idem. IV. "The Simultaneous Formation of Ortho-, Meta-, and Para-Substituted Derivatives of Benzene"

1. Amer. Chem. Soc. 37, 863-883 (1915) See Chapter XI

H S. Fry . Idem V "A Reply to A. F. Holleman," J.

K G. Falk and J. M Nelson: "Electron Conception of Valence VII. Theory of Electrolytic Dissociation and Chemical Action." J. Amer. Chem. Soc., 37, 1732-1748 (1915). The authors hold that changes occurring in chemical reactions do not depend upon the electrolytic dissociation of the reacting substances. The chemical changes are accompanied very often by electrolytic dissociation phenomena but the latter parallel the former, or vice versa, and do not necessarily piecede or cause them.

H S F1y. "Einige Anwendungen det elektronischen Auffassung positiver und negativet Valenzen VI Über die Existenz und die Eigenschaften 'fieier Radikale'." Zeitschr. physikal. Chem, 90, 4;8-480 (1915). See Chaptes XVI.

H. S. Fry: "Interpretations of Some Stereochemical Problems in Terms of the Electronic Conception of Positive and Negative Valences. VI Further Evidence for the Electronic Formula of Benzene and the Substitution Rule" J. Amer. Chem. Soc., 38, 1323-1327 (1916). See Chapter VIII.

H S. Fry Idem. VII. "The Action of Sodium Methylate upon the Products of Nikration of Ortho-, Pata-, and Meta-Chlorotoluenes" J Amer Chem Soc., 38, 1327-1333 (1916) See Chapter XII.

H. S. Fry: *Idem.* VIII "Further Evidence for the Electronic Tautomerism of Benzene Derivatives," *J. Amer. Chem Soc.*, 38, 1333-1338 (1916). See Chapter X.

- H. S. Fiy; Idem. IX^{*} "The Electronic Formula of Benzene and the Molecular Volumes of the Chlorobenzenes" f Amer. Chem. Sec, 39, 1688-1699 (1917) See Chapter XVII.
- L. W. Jones: "Electromerism, A Case of Chemical Isomesian resulting from a Difference in Distribution of Valence Electrons". Science, Vol XLVI., No 1195 (1917) The essential features of this paper have been reviewed in the abstract of Jones' paper on "Electromers and Stereomers with Postrive and Negative Hydroxyl" (p. 277).
- M. T. Hanke and K. K. Koessler, "The Electronic Constrution of Acetoacetic and Citric Acids and some of their Derivatives" J. Amer Chem. Soc., 40, 1726-1732 (1918) When citric acid is treated with fuming sulphuric acid the central carboxyl group escapes as carbon monoxide. The reason for the elimination of the central carboxyl group and the nonelimination of the two end carboxyl groups is due to a difference in the direction of the valence force holding these groups. In other words, as F1v has shown, the elimination of a carboxvl group as carbon monoxide signifies that said group functions negatively Its carbon atom corresponds to the state of oxidation and the electronic type of the carbon atom in formic acid, which readily yields carbon monoxide. Citric acid, losing carbon monoxide, yields acetone dicarbonic acid which contains the two end carboxyl groups of the original citue acid. These are eliminated as carbon droxide on treatment with caustic alkali solution, yielding acetone Again, as Fry has shown, the elimination of a carboxyl group as carbon droxide shows that said carboxyl group functions positively Its carbon atom corresponds to the state of oxidation and to the electronic type of the carbon atom in carbonic acid, which loses carbon dioxide By the application of these pre-ciples, Hanke and Koessler are able to propose electronic formulæ for various complex acids which are readily correlated with their chemical properties

Rajendralal De. "Polar and Non-Polar Valency". Trans. Chem Soc., 115, 127-134 (1919) In the light of recent theories of atomic structine, De has discussed the structural significance of complex salts such as the cobalt- and platinum-ammonia compounds, acetylides, etc., in relation to polar and non-polar valency.

Eustace J. Cuy; "The Electronic Constitution of Normal

Carbon Chain Compounds, Saturated and Unsaturated". J. Amer Chem Sac, 42, 503-514 (1920) Cuy summarizes his paper thus. "Assuming that carbon compounds are polar in nature and that carbon atoms in a chart tend to assume alternately positive and negative changes, it has been shown that the fluctuation in the various physical properties of these compounds such as melting points, boiling points, and so forth, between the even and the odd members of a given series, may be accounted for. Various reactions in which these compounds take part, such as the addition of halogen acids and the isomeric rearrangement of the halides may likewise be accounted for, on the basis of these assumotions."

Many significant papers have not been noted in the foregoing review for the reason that they are concerned chiefly with the problem of the constitution of the atom. While they bear more or less directly upon the question of chemical valence, they are not primarily direct applications of the electronic conception of valence to the interpretation of specific chemical reactions. In this connection the following papers should be noted '—

- G. N. Lewis, "The Atom and the Molecule" J. Amer. Chem. Soc., 38, 762-784 (1916)
- G N. Lews "Steric Hindiance and the Existence of Odd Molecules (Fice Radicals)". Proc Nat Acad. Sci., 2, 586 592 (1916)

Irving Langmuit "The Arrangement of Electrons in Atoms and Molecules" J. Amer. Chem. Soc., 41, 868-934 (1919).

Irving Langmuir. "Isomorphism, Isostetism, and Covalence" J. Amer Chem. Soc. 41, 1543-1559 (1019)

Irving Langmuir "The Structure of Atoms and Its Bearing on Chemical Valence". J Inc. Eng. Chem., 12, 386-388 (1920).

The atomic configurations and methods of representing electronic valences presented in these and many other papers are so complicated that they do not lend themselves readily to a convenient form of graphic or structural illustration when one attempts to make application to specific chemical reactions. In other words, their adaptability to alucidate structural formula: and chemical reactions is questionable.

B. Review of Criticisms.

The introduction of the electronic conception of valence and its extension to the interpretation of chemical phenomena has met with some criticism. The papers embodying these criticisms should be noted briefly in this bibliographical review.

- P. de Heen: "The Electron Theory and Conception of Valence" Bull. Acad. Rep. Belg., 1913, 667-679. The author's chief criticism is that the electron theory does not funnsh any satisfactory account of the behaviour of the elements of variable valency. To this it may be replied that when the valence of an element is (n) that element may function electronically in (n + 1) ways. If the ordinary valence varies, the same rule applies as well to the one value for (n) as to the other value or valence (n) Numerous illustrations of this rule have been given in the monograph. Jones (loc. cit.) has indicated methods of considering electronically the nitrogen atom which may function either as a tervalent or a quinquevalent element.
- S J Bates: "The Electron Conception of Valence". J Anne. Chem Soc., 36, 789-793 (1914). Bates maintains that "on the whole the phenomena of physics are opposed to the view that in the molecule the atoms are charged with respect to one another, and to the theory of valence developed on this assumption. Chemistry contributes the most satisfactory evidence in its avour." This type of criticism is inelevant for two reasons (1) The isolated conditions under which the quoted physical phenomena are effected (e.g., light vacua, influence of positive rays, etc.) are not comparable with the conditions under which the great majority of chemical reactions take place. Therefore, both the results and conclusions are bound to differ. (2) The electronic conception of positive and negative valence plays the part of a formulative hypothesis in the interpretation and correlation of chemical and physico-chemical phenomena.
- A F. Holleman: "Substitution in the Benzene Nucleus",

 Amer. Chem. Soc. 36, 2495-2498 (1944). Holleman maintains that there is no hypothesis able to give a satisfactory explanation of the phenomena of substitution in the benzene nucleus. He states that Fry's electronic formula for benzene and rule for the explanation of substitution reactions cannot be accepted "because there are so many obsections to it." He

presents six "most important" specific objections which should he studied intimately by the reader. Fry has replied in detail to each of these objections, I. Amer. Chem. Soc., 37, 883-892 (1915), showing by direct quotations from Holleman's criticism and Fiv's original statements that Holleman's mability to correlate Fiv's rule with certain cases of substitution was due to Holleman's failure to apply to the principle of the electronic tautomeusm of benzene derivatives the generally accepted principle that, in any tautomeric equilibrium mixture, either one, or the other, or both tautomers (depending upon conditions) may interact with a given reagent. Thus the simultaneous formation of orthos, paras, and meta-substituted derivatives is readily explained. Furthermore those cases of substitution which Holleman regarded as opposed to Fry's rule were shown to conform to the rule and to the principle of the electronic tautomerism of benzene derivatives. An appreciation of the points at issue between Holleman and Fry necessitates a critical study of their original papers. It is of interest to add that in a private communication from Professor A. F. Holleman (Amsterdam, June 7, 1015), he writes, "I fear that you have gone too far in your explanations, and it will be necessary to review your hypotheses. though I acknowledge that there is a right nucleus in them". While Holleman apparently did not completely sense the significance of the principle of electronic tautomerism and the necessary part it plays in substitution reactions, I. B. Cohen 161 clearly states the principle in his seview of Fry's electronic theory of benzene substitution. Cohen concludes his summary with the statement that Fiy's theory, "in short, is so mobile, so adaptable, and so ingeniously applied as to explain most of the facts of substitution as well as many reactions of aromatic compounds".

R. F. Brunel: "A Citicism of the Electron Conception of Valence" J. Amer. Chem. Soc., 37, 709-722 (1915). Brunel concludes that any application of the theory involves the constant use of assumptions; that the electronic formule assigned to chemical compounds are inconsistent with any rule that can be proposed, that no single well-established case of electromerism is yet known, and that the evidence of physical experiments is at present opposed to the assumption on which it is based.

H. S. Fry: "The Electronic Conception of Positive and

Negative Valences". J. Amer. Chem. Soc., 37, 2368-2373 (1915).
 A reply to the preceding paper of Brunel.

There now remains for consideration a brief discussion of the status and functions of the electronic conception of positive and negative valence. This is the subject of the concluding chapter.

CHAPTER XXIV.

THE STATUS AND FUNCTION OF THE ELECTRONIC CONCEPTION OF POSITIVE AND NEGATIVE VALENCE.

THE final chapter of Recent Advances in Organic Chemistry Dy A. W. Stewart 162 discusses "modern formulæ and their failings". The introductory paragraph bears the following statement: "An unbiased survey of the fields covered by organic chemistry cannot fail to reveal to any critical mind the fact that our structural formulæ are becoming less and less able to cope with the strain which modern research is placing upon them it is true that for work-a-day purposes they still answer admirably, and from the point of view of teaching it is doubtful if anything better could be devised. But when we go into the matter beyond the mere surface, things are not so satisfactory as they may anonea to the superficial observer."

Examples are cited by Stewart to show that our formulae have ceased to be pure "reaction-formulae" and that they frequently mislead us if we attempt to draw general conclusions from them. On the other hand, researches in physics and physical chemistry are groung us glimpses of the "intimate structure of molecules". Therefore, a difficult task is encountered when one attempts to embody reaction-formulae and untimate structure of molecules simultaneously in one configuration. Stewart admits that progress along these lines will be slow, but it is the task of investigators who concern themselves with both physical and chemical properties to invent a special symbolism which will express their results and be free from the implications that are attached to ordinary formulae "Conservatism is inguined in most scientific minds; and the struggle which new ideas have before them is generally severe."

A. Status and Function of the Electronic Conception of Valence.

Apropos of the foregoing points of view of Stewart, what is the status and function of the electronic conception of positive and negative valence and the new types of formula—electronic formula—proposed, developed, and illustrated in this monograph?

The answer to this question is manifold.

- (1) It should be recalled that the electronic conception of positive and negative valence, as noted in the introduction (Chapter I.) is a formulative hypothesis. It is maintained that it should function as such in the interpretation and correlation of chemical and physico-chemical phenomena.
- (2) The electronic conception of positive and negative valence as herewith applied to the constituent atoms of relatively well-established structural formulæ, is, at its present status of development, neither primarily nor necessarily concerned with the question of the ultimate nature of chemical affinity and the constitution of the atom per se. Such questions have invariably led into fields wherein speculation predominates. Moreover, the numerous, varied, and divergent hypotheses relating to the ultimate constitution of the atom have failed, so far, to furnish a uniformly satisfactory valence hypothesis which will enable chemists to elucidate chemical formulas and reactions.

The most to be gleaned from any or all of these anomalous hypotheses is the early and relatively sumple suggestion of Sir J J Thomson that "if we interpret the bond" of the chemist as indicating a unit Faraday tube, connecting charged atoms in the molecule, the structural formulae of the chemist can be at once translated into the electrical theory". Thus, in the electronic theory, one end of a bond corresponds to a positive, the other to a negative charge, the charge being developed through the loss or the gain of an electron. It is this earlier view that is most readily and significantly adaptable to chemical formulae. Later views on the electronic nature of valence and the distribution of valence electrons have manifested little, if any, discernible utility in the direct and lucid interpretation of the mechanism of specific chemical reactions and the correlation of the varied phenomena of inorganic and organic chemistry.

(3) It should also be recalled that the translation of a structural formula into an electronic formula is not an autural procedure. Many illustrations have been presented showing that this translation is governed by a careful study of the phenomena of ionization and electrolysis, oxidation-reduction processes, and hydrolytic reactions
Each of these phenomena is readily

- interpreted in terms of electric charges, that is, positive and negative valences or the capacity of atoms and radicals to function positively and negatively. It is in conformity with these physical and chemical phenomena that the polarity of the valences or boods in a structural formula are indicated by plus and minus signs thus making the translation of a structural formula into an electronic formula both a physically and a chemically consistent procedure.
- (4) Many substituted derivatives of benzene when singly brought into hydrolytic reactions or subjected to further substitution, yield derivatives whose existence and properties compel us to conclude that the reaction involved more than one type of molecule of the initial benzene derivative. These molecules are not those of tautomers or desmotropes. They are electronic tautomers, or electromers, existing in tautomeric equilibrium, that is, manifesting the phenomenon of electronic tautomerism If this assumption, the principle of electronic fautomerism is not valid, then we are still unable to explain either the simultaneous formation of ortho-, meta-, and para-derivatives of benzene in any given substitution reaction or the chemical properties of the substituted derivatives formed which are the specific products of respective electromers While the isolation of the electromer of any benzene derivative has not as yet been effected, the derivatives of these electromers have been obtained and their chemical properties explained and correlated in terms of their electronic formulæ In other words, the simultaneously formed ortho-, meta-, and para-substituted derivatives of a given compound are the immediate and direct reaction products of the electromers of that ompound
- (5) The electronic conception of positive and negative valence, as herewith presented, developed and applied, is a formulative hypothesis. Such an hypothesis to be efficient must be inductively true at the time of its promulgation; that is, it should explain the phenomena and laws which classification has brought together in a particular branch of science. It should also be applicable to future discoveries; and, finally, it must be clashicitively suggestive in indicating lines of future is escarch. The electronic conception of positive and negative valence has, in a measure, met the first requirement in affording interpretations of the mechanism of many hitherto, unexplained chemical reactions.

Notably substitution reactions in the benzene nucleus and the anomalous behaviour of many derivatives of benzene. It has also afforded explanations of physico-chemical phenomena such as molecular volume relation-hips, absorption of light and fluor-exence. Does it also meet the other fundamental requirement of a good hypothesis, namelly, to suggest lines of future research?

B. Suggested Lines of Future Research.

A suggested line of research may be found in endeavous to isolate electromers, the existence of which in tautometic equilibrium is clearly substantiated. As noted, electromers of derivatives of benzene have not as yet been individually isolated, but a well-established case of electromersim has been pointed out by Jones (loc cit) in his electronic explanation of certain isometic derivatives of hydroxylamine originally prepared by Mcienholmer. These tepresent the first known instances of independently existing electromers, that is, compounds completely identical in structural formulæ but differing in physical and chemical properties by virtue solely of a different arrangement of the respective positive and negative valences (valence electrons) of certain constituent atoms. The isolation of other electromers depends upon future research.

Another line of suggested issearch may be located in the well-ploughed field of unsuccessful attempts to effect a direct asymmetric synthesis, i.e, to piepaie a compound which displays optical activity through rotation of the plane of polarized light by vitue of its containing an asymmetric atom, e.g., the carbon atom, united to four chemically different atoms or radicals. A suggestion to this end may exist in the following explanation of magnetic optical activity in terms of the electronic conception of positive and negative valence.

A naturally optically active carbon compound is either dextroor lavo-rotatory independently of the direction in which the plane polarized light passes through its solution. On the other hand, a magnetically optically active liquid is dextro-or lavo-rotatory according to the direction in which the light passes through it, the magnetic field being constant. Now natural optical activity of a carbon compound depends upon the presence of an asymmetric carbon atom in its molecule. Briefly illustrated (C a, b, c, a) indicates that the quadrivalent carbon atom is united to four chemically different atoms or radicals a, b, c, and d The compound (C a, b, c, c) is not optically active since two of the four atoms united to it (c and c) are chemically identical. (C a, b, c, c) displays optical activity in a magnetic field. One natural conclusion is that the two chemically identical atoms (c and c) must function differently in the magnetic field. In other words, in terms of the electronic conception, whether (c) and (c) are naturally both positive or naturally both negative, the magnetic field induces a rearrangement of the valence electrons in such a way that one (c) functions positively and the other (c) functions negatively It has been shown that a given atom may function sometimes negatively, sometimes positively, and these two states are entirely different chemically. Hence, the magnetic field may induce magnetic optical activity by converting mactive

(C a, b, c, c) or inactive (C a, b, c, c) into magnetically optically active (C a, b, c, c). This may be termed electromeric asymmetry, the occasion of magnetic optical activity.

It is also quite conceivable that a compound such as

(C a, b, c, c) in which like atoms (c and c) possess opposite polarity would not show optical activity unless in a magnetic field. In this case the magnetic field would not be the occasion of atoms (c and c) functioning positively and negatively, but it would differentiate them in their relative spatial positions in the molecule so that the carbon atom (C) is asymmetric. It displays electrometic asymmetry. This differentiation in space and in

polarity of the atoms (c and c) in a magnetic field is readily correlated with the fact that a magnetically optically active liquid is dextio- or lawo-rotatory according to the direction in which the light passes through it.

This proposed hypothesis also correlates the well-established theory of the asymmetric carbon atom of Le Bel and Van't Hoff with Faraday's discovery of and Sii W. H. Perkin's remarkable researches in the field of magnetic optical activity.

Immediately in this connection it should also be noted that many crystalline substances, such as quartz, are optically active. but it has been held by physical chemists that the activity here is not due to the arrangement of atoms within the molecule but rather to a certain undefined arrangement of crystalline particles. Now X-ray and crystal structure investigations are clearly indicating the types of the airangements of the atoms in spaced lattices. Accordingly, it is quite possible that many such arrangements present examples of electromeric asymmetry. In other words, optical activity of crystals is assumed to be due to electromeric asymmetry which exists naturally in optically active crystals but is induced in nactive substances by the magnetic field.

The suggestion of the conception "electromenic asymmetry."

do not of course, in itself convittute an explanation; but it may serve as a nucleus for the development of a more comprehensive hypothesis which should embrace and correlate, along the lines indicated, all types of optical activity. The underlying concepts are the electronic conception of positive and negative valence and the principle of asymmetry, which latter has played a vital part in the explanation of the optical activity of compounds of carbon.

The phenomena of magnetic optical activity and the proposed electionic interpretation also suggest other lines of investigation. For example, may not a sufficiently powerful magnetic field have some effect upon the relative concentrations or relative reactivities of the electromers in a tautomeric equilibrium mixture such as $(c_{\rm opt}^{\dagger}, L_{\rm opt}^{\dagger}, E_{\rm opt}^{\dagger}, E_{\rm opt}^{\dagger})$. Since benzene derivatives display electronic tautomerism, they may be expected to react

differently within a magnetic field than they do in a non-magnetic field. This difference in reactivity under, the two conditions could be ascertained by a quantitative determination of the reaction products which are the derivatives of the respective electromers. If the magnetic field altered in any way the

concentrations or reactivities of the electromers, $C_s \overset{+}{H}_s$. $\overset{+}{X}$ and $C_s\overset{+}{H}_s$. $\overset{+}{X}$, variations in the quantities of the substituted derivatives

of C_6H_5 . X and C_8H_5 . X would establish the fact.

In this connection studies of the hydrolytic reactions of optically active and inactive compounds, within and without the magnetic field, are suggested. Also, addition reactions present subjects for investigation. For example, the addition of the halogens and halogen acids to unsaturated compounds have special significance in attempts to effect asymmetric synthesis.

The two types of double bonds, continuous and diview, proposed by the author afford a ready explanation of the varied and apparently anomalous addition reactions frequently noted in the literature. How will a magnetic field effect these addition reactions?

The variable physical conditions under which these suggested reactions may be conducted must not be overlooked. Electronic fautomers are undoubtedly very subtle. The behaviour of electromers and the possibility of their isolation are quite likely to be influenced by the magnetic field, its character and intensity; by the solvent medium, its density, molecular volume, temperature, dielectric constant, chemical properties, etc; and by the presence of catalysts, either directly employed or as secondary products of the reaction. All of these conditions as well as the quantities of the reactions must be accurately standardized and carefully determined in any of the surgested iesearches.

From the foregoing it is quite apparent that the electionic conception of positive and negative valence meets the two fundamental requirements of an efficient formulative hypothesis: it not only explains many facts and phenomena—its first function—but it is also deductively suggestive in that it has indicated various lines of future research. These hypotheses are proposed with the understanding that assumption is not necessarily presumption.

Finally, a statement made in the introductory chapter will now bear repetition: "In view of the fact that electronic formulae, in many instances, have proven to be more precise and more significant than the customary structural formulae in the explanation of chemical and physico-chemical phenomena and the mechanism of reactions, it is quite conceivable that the electronic conception of positive and negative valence as a formulative hypothesis may become a necessary adjunct to the structure theory. This, of course, must depend upon the nature and extent of its applications and experimental venifications, and upon the part that should be played by just criticisms in bringing to light the relative merits and demerits of its applications." Complementary to this opinion the author begs to conclude this monograph by quoting the final paragraph of A. W. Stewart's Recent Advances in Organue Chemistry (to, cit.)—

"It is not to be expected that success will be attained at a stroke. Much more probably, there will be a good deal of fumbling and recesting to be gone through, just as there was before our present-day formulae emerged from the melting-pot. Any suggestions, therefore, which tend towards the enlargement of our ideas of chemical constitution should be welcomed by those who have sufficient critical spirit to guasp the failure of our contemporary formulae under the stain of modern investigations."